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(54) Title: TRANSGENIC ORGANISM

(57) Abstract

A transgenic organism is described which has an increased starch yield. The preferred embodiment concerns a plant or plant cell containing a recombinant DNA construct containing, in operational relationship to a plant promoter sequence or sequences enabling the expression of the gene by the plant or plant cell thereby to enhance the rate of production and/or yield of starch by the plant or the plant cell, a DNA sequence encoding an exogenous ADP glucose pyrophosphorylase enzyme (AGP) or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme, characterised in that the said DNA sequence is the gene sequence, including non-critical allelic variations thereof, encoding the barley Hordeum vulgare endosperm AGP or an active sub-unit thereof, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units.

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#### TRANSGENIC ORGANISM

The present invention relates to a transgenic organism.

In particular, the present invention relates to a transgenic starch producing organism having an increased ability to synthesize starch and one that is capable of producing starch in high yields. More particularly the present invention relates to a transgenic organism comprising a nucleotide sequence coding for exogenous ADP glucose pyrophosphorylase (AGP).

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In a preferred embodiment the present invention relates to a transgenic plant or plant cell capable of expressing exogenous AGP in the starch producing centres in the plant, namely the chloroplasts and the amyloplasts. The invention also relates to a recombinant DNA construct for use in the transformation of a plants or plant cell showing enhanced starch production, and plants and plant cells transformed with the recombinant DNA construct.

ADP glucose pyrophosphorylase (E.C.2.7.7.27) (AGP) is one of the primary enzymes involved in the biosynthesis of starch and glycogen in organisms such as plants, algae, fungi and bacteria, particularly plants.

AGP catalyses the reaction:

$$\alpha$$
-glucose-1-P + ATP  $\leftarrow$  ADP-glucose + PP<sub>1</sub>

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the product ADP-glucose being the major donor of glucose in the biosynthesis of starch in plants. Moreover, that reaction has been shown to be the rate limiting factor in the synthesis of starch in organisms such as plants, the rate of that reaction in turn being critically dependent upon the AGP concentration. Because of that, AGP has been the subject of intensive investigation and for a general review of recent studies on plant AGP, reference should be made to Kleczkowski et al: Z. Naturforsch. 46c, 605-612 (1991).

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As reported by Kleczkowski et al (*ibid*) and elsewhere, AGP is widely distributed throughout the plant kingdom and is found in some starch producing bacteria, such as E. coli. Plant AGP exists as a tetramer (210 to 240 kDa) composed of two small sub-units (50 to 55 kDa) and two large sub-units (51 to 60 kDa) in contrast to bacterial AGP which appears to consist of four units of equal size. AGP has also been shown to be produced in cyanobacteria and in algae, where its tetrameric structure is similar to that in plants, i.e. two large and two small sub-units, rather than the homotetrameric structure found in ordinary bacteria.

Because of the commercial importance of starch, primarily as a foodstuff but also as an important industrial chemical, AGP itself and recombinant DNA constructs containing DNA sequences encoding AGP for the transfection of plants and plant cells as a means of increasing plant AGP concentration and hence increased biosynthesis of starch in plants and increased starch yields, have formed the subject matter of several recently published patent applications.

For example, in EP-A-0368506 a method of extracting AGP from wheat leaf and wheat endosperm is disclosed. Also disclosed are the cDNA sequences encoding wheat leaf and wheat endosperm AGP, and various plasmids containing those sequences for subsequent insertion into plants to provide plants having an increased ability to synthesise starch, although that latter step is not described in detail, nor are any examples given of transgenic plants containing those constructs.

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WO 91/19806 discloses transformed plant cells and plants having elevated levels of starch and starch biosynthesis achieved by incorporating into the plant genome a DNA construct comprising in sequence a plant promoter, a DNA sequence encoding a fusion polypeptide consisting of a plastid transit peptide and a bacterial (E. coli) AGP, and a 3'-non-translated region which functions in the plant cell to cause transcriptional termination and the addition of a polyadenylated tail to the 3'- end of the corresponding DNA sequence. The DNA sequence encoding E. coli AGP is given, as well as the deduced amino acid sequence. Transgenic potato and tomato plants transformed with the E. coli AGP gene are shown to produce increased starch

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yields. It is suggested that other bacterial sources besides <u>E. coli</u>, and also algae, may be used as a source for the AGP gene to be used in the transformation of the plants and plant cells to provide increased starch yields. However, there is no mention of the isolation of nucleotide sequences coding for AGP enzymes from those other sources or their expression in such transgenic systems.

A similar disclosure is contained in WO 92/11382 which likewise discloses the transformation of plants, especially potato plants, with bacterial (E. coli) DNA encoding bacterial AGP, with the objective of increasing starch biosynthesis and starch yield in such plants.

A slightly different objective is set out in EP-A-0455316. There the objective is to increase sugar and protein concentrations in plant-based foodstuffs at the expense of starch formation. That is achieved by incorporating into the plant genome a DNA sequence encoding AGP, but in an inverted orientation in the transformation vector. Transcription of the reversed sequence results in an anti-sense mRNA which inhibits the production of AGP in the plant cell leading to reduced AGP activity and reduced starch production.

All plant AGPs investigated so far have been reported to be strongly activated by 3-phosphoglycerate (PGA) and inhibited by inorganic phosphate (P<sub>i</sub>). Also, the PGA/P<sub>i</sub> ratio in the chloroplasts and amyloplasts where biosynthetic starch production is concentrated is believed to play a key regulatory role in starch synthesis. It is known, for example, that chloroplast PGA/P<sub>i</sub> ratios are at the highest activity during the daylight hours, i.e. during photosynthesis, which period coincides with the peak period of starch production in the chloroplasts. The regulation of the AGP formation in non-photosynthetic tissues is less well understood, but the activatory and inhibitory roles of PGA and P<sub>i</sub>, respectively, i.e. the PGA/P<sub>i</sub> ratio, is believed still to play an important part.

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The present invention addresses the problem of how to increase AGPase levels and/or starch levels in starch producing organisms.

According to a first aspect of the present invention there is provided a transgenic starch producing organism comprising a nucleotide sequence coding for an exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme, wherein the nucleotide sequence is capable of being expressed in the organism; characterised in that the activity of the enzyme or sub-unit thereof is substantially independent of any level of *in vivo* 3-phosphoglycerate and/or any *in vivo* level of inorganic phosphate; and further characterised in that the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

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According to a second aspect of the present invention there is provided a transgenic starch producing organism comprising exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme, wherein the activity of the enzyme or sub-unit thereof is substantially independent of any level of *in vivo* 3-phospho-glycerate and/or any *in vivo* level of inorganic phosphate and wherein the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

According to a third aspect of the present invention there is provided a potato tuber containing an enhanced starch content.

According to a fourth aspect of the present invention there is provided a method of increasing the rate and/or yield of starch production in an organism, especially a plant or a plant cell, which method comprises introducing into an organism a nucleotide sequence according to the present invention to form a transgenic organism according to the present invention and expressing the nucleotide sequence.

According to a fifth aspect of the present invention there is provided a method of increasing the rate and/or yield of starch production in an organism, especially a plant or a plant cell, which method comprises introducing into or forming in an organism a ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof according to the present invention.

According to a sixth aspect of the present invention there is provided any one of the following: A cDNA sequence identified herein as SEQ ID No. 2, including non-critical allelic variations of that sequence; An amino acid sequence as shown in SEQ ID No. 4, including variants thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence; A cDNA sequence identified herein as SEQ ID No. 5 including non-critical allelic variations of that sequence; An amino acid sequence as shown in SEQ ID No. 6, including variants thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.

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According to a seventh aspect of the present invention there is provided any one of the following plasmids: Plasmid pPPS1; Plasmid pPPL1; Plasmid pPPL1M; Plasmid pPPL4; Plasmid pPPL4; Plasmid pPPL4; Plasmid pPPL5; Plasmid pBKL4; Plasmid pVictor IV GN.

According to an eighth aspect of the present invention there is provided a method of 15 increasing the rate and/or yield of starch production in an organism, especially a plant or a plant cell, which method comprises introducing into an organism a recombinant DNA construct containing an exogenous DNA sequence encoding an exogenous ADP glucose pyrophosphorylase enzyme (AGP) or sub-unit thereof and one or more 20 promoter sequences enabling the expression of the AGP encoded by that sequence by the organism thereby to increase the AGP content of the organism and in consequence to increase the rate of starch production by the organism and/or the starch yield. characterised in that the said DNA sequence is the gene sequence encoding the barley (Hordeum vulgare) endosperm AGP or a sub-unit thereof, or a variant thereof having 25 non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units. wherein the construct is expressed in the organism; characterised in that the activity of the enzyme or sub-unit thereof is substantially independent of any level of in vivo 3-phospho-glycerate and/or any in vivo level of inorganic phosphate; and further 30 characterised in that the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

According to a ninth aspect of the present invention there is provided a vector for the transformation of an organism, especially a plant or a plant cell, to increase the AGP content of such an organism consequently to increase the rate of starch production by such an organism, such vector comprising a recombinant DNA construct containing a DNA sequence encoding an exogenous ADP glucose pyrophosphorylase enzyme (AGP), such vector also incorporating the necessary promoter and other sequences enabling the expression of that exogenous AGP in an organism transformed by that vector, characterised in that the said DNA sequence is the gene sequence encoding the barley (Hordeum vulgare) endosperm AGP or a sub-unit thereof, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units, wherein the construct is capable of being expressed in the organism; characterised in that the activity of the enzyme or sub-unit thereof is substantially independent of any level of in vivo 3-phospho-glycerate and/or any in vivo level of inorganic phosphate; and further characterised in that the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

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According to a tenth aspect of the present invention there is provided a method of targeting an exogenous protein to the amyloplast of plants or plant cells which comprises introducing into the plant or plant cell a recombinant DNA construct containing a DNA sequence encoding a starch branching enzyme transit peptide and an exogenous DNA sequence encoding the exogenous protein; wherein the construct is capable of being expressed in the plant or plant cells; preferably wherein the DNA sequence encoding the starch branching enzyme comprises the sequence identified as SEQ.ID.No.5 and/or the starch branching enzyme expressed in the plant or plant cell by said construct comprises the amino acid sequence identified as SEQ.ID. No. 6.

According to an eleventh aspect of the present invention there is provided an AGP enzyme or sub-unit thereof whose *in vivo* activity is substantially independent of any level of *in vivo* 3-phospho-glycerate and/or any *in vivo* level of P<sub>i</sub>, and whose activity is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

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According to a twelfth aspect of the present invention there is provided a foodstuff made from or comprising an organism according to the present invention; preferably wherein the foodstuff is a fried foodstuff; more preferably wherein the foodstuff is a potato.

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The term 'transgenic organism' in relation to the present invention means an organism comprising an expressable exogeneous nucleotide sequence or an expressed product of such an expressable exogeneous nucleotide sequence. Preferably the expressable exogeneous nucleotide sequence is incorporated in the genome of the organism.

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The term 'organism' in relation to the present invention includes any starch producing organisms such as plants, algae, fungi and bacteria, as well as cells thereof. Preferably the term means a plant or cell thereof, more preferably a potato and especially a potato tuber.

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The term 'nucleotide' in relation to the present invention includes genomic DNA, cDNA, synthetic DNA, and RNA. Preferably it means DNA, more preferably cDNA.

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The terms 'allele' and 'variant' in relation to the present invention mean any substitution of, variation of, modification of, replacement of, deletion of or the addition of one or more nucleic acid(s)/amino acids from or to the sequence providing the resultant sequence expresses or exhibits the required enzymatic activity. They also mean a substantial homologous sequence wherein there is homology with respect to at least the essential nucleic acids/nucleic acid residues/amino acids for expression of or exhibition of the required enzymatic activity. Preferably there is at least 80% homology, more preferably at least 90% homology, and even more preferably there is at least 95% homology with the listed sequences. Hybrid sequences are also covered. These may be prepared from at least two different sources - e.g. the variant may include a sequence from one source that gives the variant the independence vis-a-vis the level of in vivo 3-phospho-glycerate and a sequence from another source that gives the variant the independence vis-a-vis the level of in vivo P<sub>i</sub>.

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The term 'sub-unit' in relation to the present invention means an active component of the enzyme that displays the required enzymatic activity. For example, in the case of AGP enzyme obtained from barley (<u>Hordeum vulgare</u>), which is a heterotetramer containing two large sub-units and two small sub-units, the term includes any one of those sub-units as well as combinations thereof as well as a shortened variant thereof.

The term 'retains enzymatic activity' in relation to the present invention means that the activity of the enzyme variant or sub-unit thereof is still substantially independent of any level of *in vivo* 3-phospho-glycerate and/or any *in vivo* level of inorganic phosphate, but not necessarily to the same extent as the native enzyme.

The term 'substantially independent' in relation to the present invention means that the enzyme has a decreased sensitivity to levels of PGA and/or of inorganic phosphate, preferably at least to PGA. By way of example, in the absence of PGA the levels of AGPase activity of the native enzyme or sub-unit thereof are in the order of at least 0.002 units per mg protein, preferably at least 0.01 units per mg protein when measured in *Bintje* potato tuber extract. Typically, in the case of the preferred barley AGP enzyme we have found that the AGPase levels of the large sub-unit are greater than the levels of the small sub-unit and are typically in the order of greater than 0.02 units per mg protein and can be in the order of 0.05 units per mg protein when measured in *Bintje* potato tuber extract. This is in contrast to the known enzymes which have no, or at most negligible, AGPase activity in the absence of PGA.

25 Preferably the enzymatic activity of the AGP enzyme is at least substantially independent of any level of *in vivo* 3-phospho-glycerate.

More preferably the enzymatic activity of the AGP enzyme is not stimulated by fructose-1,6-bisP and it is not inhibited by AMP.

Preferably the AGP enzyme is a heteromer, preferably a heterotetramer, more preferably a heteromer containing two large sub-units and two small sub-units.

Preferably the AGP enzyme is isolatable from <u>Hordeum</u>, preferably wherein the enzyme is barley (<u>Hordeum vulgare</u>) endosperm AGP or a sub-unit thereof, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units.

Preferably the nucleotide sequence is a DNA sequence.

Preferably the DNA sequence encodes the large sub-unit of the barley endosperm

AGP or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequence defining the large sub-unit of the barley endosperm AGP.

Preferably the DNA sequence is the sequence identified herein as SEQ ID No 1, including non-critical allelic variations of that sequence.

Preferably the DNA sequence encodes the small sub-unit of the barley endosperm AGP, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequence defining the small sub-unit of the barley endosperm AGP.

Preferably the DNA sequence is the sequence identified herein as SEQ ID No 2, including non-critical allelic variations of that sequence.

25 Preferably both of the DNA sequences are expressed in the same organism. The DNA sequences need not be derived from the same initial source, such as barley. It is preferred however that they are from the same source, for example barley.

Preferably, when both of the DNA sequences are expressed in the same organism,

each DNA sequence addditionally codes for a different marker - e.g. the large or
small sub-unit of barley AGP enzyme may be in a construct that contains a kanamycin
resistance gene such as a construct based on plasmid pBKL4 or pVictor IV GN and

another small or large sub-unit of barley AGP enzyme may be in a construct that contains a mannose isomerase gene such as a construct based on plasmid pVictor IV SGiN Man.

Preferably the expressed AGP enzyme or sub-unit thereof comprises the amino acid sequence set out in SEQ ID No. 3, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.

Preferably the expressed AGP enzyme or sub-unit thereof comprises the amino acid sequence set out in SEQ ID No. 4, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.

Preferably the expressed AGP comprises both a large sub-unit having the amino acid sequence set out in SEQ ID No. 3 or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence, and a small sub-unit having the amino acid sequence set out in SEQ ID No. 4 or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.

Preferably the nucleotide sequence additionally codes for a transit peptide which can transport, or assist in the transportation of, the enzyme or sub-unit thereof from the cytoplasm to the relevant or appropriate plastid(s), such as a chloroplast and/or an amyloplast. Preferably the transit peptide is Rubisco Activase transit peptide or Starch Branching enzyme transit peptide.

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Preferably the transit peptide is coded for by a DNA sequence comprising the sequence identified as SEQ. I.D. No. 5, including non-critical allelic variations of that sequence.

Preferably the transit peptide has an amino acid sequence comprising the sequence identified as Seq.I.D.No. 6, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.

Preferably the nucleotide sequence is operatively connected to a promoter which expresses the sequence wherein the promoter is cell, tissue or organ specific.

Preferably the promoter has the sequence identified as SEQ.I.D.No.7, or a variant thereof having non-critical nucleotide substitution(s) or deletion(s) at one or more locations in that sequence.

Preferably the AGP enzyme or sub-unit thereof comprises the amino acid sequence set out in SEQ ID No. 3, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence and/or the amino acid sequence set out in SEQ ID No. 4, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.

Preferably the organism is a transgenic plant.

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Preferably the transgenic plant is a potato plant.

Preferably the nucleotide sequence according to the present invention is obtainable from any one of the following plasmids: Plasmid pVictor IV SGiN Man; Plasmid pPPS1; Plasmid pPPL1; Plasmid pPPL1M; Plasmid pPPS4; Plasmid pPPL4; Plasmid pPPL5; Plasmid pBKL4; Plasmid pVictor IV GN.

Preferably the enzyme is obtainable from a eukaryotic source.

The present invention has broad applicability to starch producing organisms, especially plants. The present invention works better in organisms such as plants compared to bacteria.

In particular the present invention works better in plants compared to <u>E. Coli</u> where 30 AGP activity is stimulated by fructose-1,6-bisP and inhibited by AMP. This <u>E. Coli</u> pathway is different to the pathway for the biosynthesis of starch in plants and algae.

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With regard to one preferred aspect of the present invention, namely a foodstuff prepared from frying a potato according to the present invention, it is to be noted that the increased starch content of the potato will lead to less fat/oil uptake during frying. This results in obvious dietary advantages. Moreover, the increased levels of starch also means that there are decreased free levels of reducing carbohydrates - which are used in starch synthesis - and so there is a decreased tendancy for the resultant product to become discoloured on frying by example reaction of the reducing carbohydrates with the hot fat/oil.

In accordance with a preferred aspect of the present invention it was found that AGP from barley (Hordeum vulgare) endosperm is highly active even in the absence of the activator PGA and is relatively insensitive to PGA/P<sub>i</sub> ratios which play an important regulatory function in the case of AGP from most other known plant sources. The PGA/P<sub>i</sub> ratio is also believed to play an important regulatory function in non-plant AGP, e.g. algal AGP.

The cDNA sequences encoding parts of the large and the small sub-units of the barley endosperm AGP and the deduced amino acid sequences have recently been established and published in Plant Molecular Biology, 19, 381-389 (1992). The complete DNA sequence encoding the large sub-unit together with the cDNA for the large sub-unit are set out in Plant Physiol. 100, 1617-1618, (1992).

In accordance with the present invention the complete DNA sequence encoding the small sub-unit of the barley endosperm AGP and the deduced amino acid sequence has now been established. Those complete cDNA sequences are reproduced herein as SEQ ID Nos 1 and 2 encoding, respectively, the large and small sub-units of the barley endosperm AGP, whilst the deduced amino acid sequences are set out herein as SEQ ID Nos 3 and 4, respectively.

Thus, in the preferred embodiment of the present invention, it was discovered that starch production in plants can be enhanced/increased by incorporating into the plant's genome and under the control of suitable promoter sequence or sequences

promoting the expression of the gene in the plant cells, particularly in the chloroplasts and amyloplasts, DNA sequences encoding either the large (60 kDa) sub-unit of barley (Hordeum vulgare) endosperm AGP or the small (51 kDa) unit, or both.

Thus, in a highly preferred aspect of the present invention there is provided transgenic plants and plant cells having increased rates of starch production and/or starch content, as compared with the corresponding non-transformed plant or plant cell, such plants and plant cells having been transformed with a recombinant DNA construct containing, in operational relationship (particularly in downstream relationship) to a plant promoter sequence or sequences enabling the expression of the gene in the plant or plant cell, the gene sequence encoding the barley (Hordeum vulgare) endosperm AGP or an active sub-unit thereof retaining the enzymatic activity of the heterotetrameric AGP, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in the amino acid sequences defining the barley endosperm AGP or either of its sub-units.

Whilst, in accordance with the present invention, a wide variety of organisms (e.g. plants and plant cells) may be transformed (especially with the gene encoding barley endosperm AGP or either of its sub-units) to increase starch production and starch yields in that particular organism, the preferred embodiment concerns the transformation of the major starch producing plant crops, namely potato, rice, wheat and maize, which four crops in terms of calorific value, probably account for three quarters of the world's food supply. Sugar beet may also be transformed.

In a more specific aspect of the present invention there are provided transgenic plants and plant cells having increased rates of starch production and/or providing increased starch yields compared with the non-transformed material, such plants and plant cells having been transformed with a recombinant DNA construct containing in downstream relationship to a plant promoter sequence or sequences enabling the expression of the gene in the transformed plant or plant cells, either or both the sequences SEQ ID No 1 and SEQ ID No 2 as set out in the prescribed fashion in the sequence listings annexed hereto and which are taken to be part of the present

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specification, or an allelic variant of either sequence showing substantial homology with the listed sequence and containing non-critical nucleotide substitutions at one or more locations in the nucleotide chain.

Alternatively defined, there are provided, in accordance with the present invention, transgenic plants and plant cells showing enhanced levels of AGP production, particularly, in the chloroplasts and amyloplasts, such plants and plant cells having been transformed with a recombinant DNA construct enabling the expression within the plant or plant cells of barley endosperm AGP or either of its sub-units, those sub-units having the derived amino acid sequences set out in SEQ ID Nos. 3 and 4, or a variant of such a sequence having non-critical amino acid substitution(s) or deletion(s) at one or more locations in the amino acid sequence defining the barley endosperm AGP or either of its sub-units.

15 Also provided in accordance with this invention are plant transformation vectors for the transformation of plants and plant cells to increase the AGP content of such plants and plant cells and thus to increase the rate of production of starch by the transformed plant or plant cell and/or the starch yield, such vectors containing one or more promoter sequences functional in plants linked in operational relationship with a DNA sequence encoding barley endosperm AGP, or either of its sub-units. More especially plant transformation vectors are provided comprising one or more promoter sequences functional in plants linked in operational relationship with either or both the sequences SEQ ID No 1 or SEQ ID No 2, or an allelic variant of either sequence showing substantial (at least 80%) homology with the listed sequence but having non-critical nucleotide substitution(s) at one or more locations in the nucleotide chain.

With regard to the promoter, numerous promoters which are functional in plants are known. The promoter should be capable of allowing suffient expression to result in the desired increase in starch production. Preferably, the promoter should be chosen so that the increased starch production is carried out in the plant tissues where the starch production is required. For instance the promoters of starch biosynthetic genes from plants may be useful.

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Known examples of such promoters include the promoter of the granule bound starch synthase gene from potato (Van der Leij et al. [1991] Mol. Gen. Genet. 228: 240-248), and the promoter of the starch branching enzyme gene Sbe 1 from rice (Kawasaki et al. [1993] Mol. Gen. Genet. 237: 1-16).

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For expression in potato, a tuber specific class I patatin promoter is preferred (Mignery et al. [1988] Gene. 62: 27-44). The DNA sequence encoding the tuber specific class I patatin promoter is set out in the appendix hereto as SEQ ID No. 7. This patatin promoter was obtained from Dr. William Belknap, USDA - ARS, Alabany, California.

The DNA sequence encoding barley endosperm AGP is preferably linked to other control sequences for the expression of the DNA in addition to a promoter sequence such as a transcription terminator sequence. Transcription terminators may be derived from a variety of different genes, including plant, viral and Agrobacterium genes. A cauliflower mosiac virus 35S terminator is preferred.

AGP activity can occur in different sites in plants. For example in potatoes AGP activity is mainly localised in the chloroplasts (i.e. plastids specialising in photosynthesis) or the amyloplasts (i.e. plastids specialising in starch storage). Many amyloplast-localised proteins are expressed as precursors and are targeted to the amyloplast by an appropriate transit peptide that is subsequently removed. Similarly, many chloroplast-localised proteins are expressed as precursors which can be targeted to the chloroplast by an appropriate target peptide.

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Whilst not wishing to be bound by any theory, it is believed that both the large and small sub-units of the barley endosperm AGP are synthesised as precursor peptides. Additional sequences are found to be attached to the amino-termini of the mature proteins which are understood to represent transit peptides. The transit peptide is then cleaved upon sequestration of the presursor protein into the plastid. It is understood that the enzyme is not subjected to any other post-translation modification process in vivo.

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However, in another embodiment of the present invention, it is desirable for the AGP transit peptides to be supplemented with one or more additional transit peptides. The transit peptide can be conveniently fused directly at the amino terminal methionine of the AGP barley sub-unit. In further preferred embodiments the barley AGP transit peptide can be substituted by another amyloplast or chloroplast transit peptide. The barley endosperm AGP cDNA is inserted into a convenient cloning vector, e.g. a plasmid, at a suitable restriction site. The DNA sequence of interest can be encloned into further vectors, if necessary, for the incorporation of additional DNA sequences. Suitable plant transit peptides include known chloroplast (Gavel & Von Heine [1990] FEBS Lett. 261: 455-458) or amyloplast (Van der Leij et al. [1991] Mol. Genet. 228: 240-248; Klosgen et al. [1989] Mol. Gen. Genet. 217: 155-161; Brisson et al. The Plant Cell [1989] 1: 559-566) transit peptides.

In potatoes, preferably a rubisco activase transit peptide (Werneke et al. Proc. Natl. Sci. USA [1988] 85: 787-791) or a starch branching enzyme transit peptide is used. The 480 bp starch branching enzyme cDNA sequence from potato showing 120 nucleotides of the 5<sup>1</sup> untranslated region and 360 nucleotides of the coding region (see SEQ.I.D. No. 5), which contains a putative 75 amino acid transit peptide and 45 amino acids of the mature branching enzyme is set out in the appendix hereto as SEQ ID No. 6.

In addition to the transit peptide portion of a protein, it may be desirable to include sequences encoding a portion of the mature plastid-targeted protein to further facilitate intracellular transport.

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Preferably the plasmids are also provided with selection markers to enable the transformed plant cells to be separated out from plant cells which have not been transformed. Suitable genes are known and include e.g. a neomycin phosphotransferase gene (e.g. neo npt II), a phosphinotricine/bialaphos acetyl-transferase gene (e.g. bar) and a \( \beta\)-glucuronidase gene (e.g. uidA) or a phosphomannose isomerase gene (e.g. manA, pmi).

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In a preferred embodiment, the transformation vectors may be prepared by initially obtaining cDNA encoding the small and large units of barley endosperm AGP by the method described in Plant Molecular Biology, 19, 381-389 (1992). For ligation into a convenient cloning vector, e.g. a plasmid, the barley endosperm AGP cDNA is provided with restriction sites at each end by PCR using the oligonucleotide primers obtained by conventional oligonucleotide synthesis procedures or a commercially available oligonucleotide synthesizer such as, for example, Applied Biosystems 381 A DNA synthesizer. These restriction sites should be homologous with sequences in the cloning vector. The desired DNA sequence can be recloned into further vectors for preparation of the ultimate transformation vectors for preparing the transgenic starch producing organism, especially a transgenic plant.

In the preferred embodiment of the present invention, the plant or plant cells may be transformed by any suitable technique for transforming cells - such as use of T-DNA, electroporation, injection, DNA bombardment or fusion. After transformation, a whole plant can be cultivated from a transformed plant cell in the usual manner.

Preferably, transformation of the plant cell is achieved with T-DNA using Agrobacterium tumefaciens or Agrobacterium rhizogenes. If agrobacteria are used for transformation, the barley endosperm cDNA needs to be incorporated initially into either an intermediate vector or a binary vector. The intermediate vectors can be integrated into Agrobacterium tumefaciens by means of a helper plasmid. Preferably binary vectors are used, which can be transformed directly into agrobacteria. Binary vectors comprise a selection marker gene and a linker or polylinker which are framed by the right and left T-DNA border regions. The agrobacteria used as host cell should comprise a plasmid carrying a vir region, which is necessary for the transfer of the T-DNA into the plant cell. Transformation using Agrobacterium is achieved by cultivating the Agrobacterium with the plant cell.

Depending on the plant species to be transformed, a variety of different plant transformation vectors can be used. These include pBIN 121, pAL4404, pEHA101, pBKL4, pVictor IV SGiNMan and pVictor IV GN.

For the transformation of potato species Agrobacteria the preferred plant transformation vectors are the plasmids pBKL4, pVictor IV SGiNMan and pVictor IV GN. These plasmids are described later in greater detail.

Preferred plasmids used in the construction of the plasmid used for transformations include pPATA1 and pBluescript II KS. Plasmid pBluescript II KS is a widely used cloning vector available from Stratagene.

Plants can be confirmed as transformed by performing conventional blotting assays and PCR.

The starch content of the plants can be analysed based upon the specific gravity determined using the weight in water and the weight in air as described by W.A. Gould In: Chipping Potato Handbook, ed. Gould, W.A. The Snack Food Association, Vermont, 1989, pp 18-22, in an article entitled "Specific gravity, its measurement and use.

The limitation of the exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme being not stimulated by fructose-1,6-bisP and/or not inhibited by AMP, which further distinguishes the present invention from the AGP enzymes of the prior art such as those of WO 91/19806 and WO 92/11382, can be expressed in the alternative as either the exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme not being only just an E. Coli AGP enzyme, or the exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme being capable of catalysing the reaction

$$\alpha$$
-glucose-1-P + ATP  $\leftarrow$  ADP-glucose + PP<sub>1</sub>.

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The following samples were deposited in accordance with the Budapest Treaty at the recognised depositary The National Collections of Industrial and Marine Bacteria Limited (NCIMB) at 23 St. Machar Drive, Aberdeen, Scotland, United Kingdom, AB2 1RY on 29 March 1994:

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- E. Coli containing plasmid pPPS1 (NCIMB 40618);
- E. Coli containing plasmid pPPL1 (NCIMB 40619);
- 10 E. Coli containing plasmid pPPS4 (NCIMB 40620);
  - E. Coli containing plasmid pPPL4 (NCIMB 40621); and
  - E. Coli containing plasmid pPPL5 (NCIMB 40622).

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The following samples were deposited in accordance with the Budapest Treaty at the recognised depositary The National Collections of Industrial and Marine Bacteria Limited (NCIMB) at 23 St. Machar Drive, Aberdeen, Scotland, United Kingdom, AB2 1RY on 31 March 1994:

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- E. Coli containing plasmid pBKL4 (NCIMB 40623);
- E. Coli containing plasmid pVictor IV GN (NCIMB 40624); and
- 25 E. Coli containing plasmid pVictor IV SGiN Man (NCIMB 40625).

A detailed construction of plant transformation vectors according to the present invention and the transformation of plants and plant cells using those vectors to produce transgenic plants according to this invention having increased rates of starch biosynthesis and/or starch yield will now be described in more detail.

In this regard, the present invention will now be described only by way of example.

In the following Examples reference is made to the accompanying figures in which:

- 5 Figure 1 shows the restriction map for plasmid pPATA1;
  - Figure 2 shows the restriction map for plant transformation vector pBKL4;
  - Figure 3 shows the restriction map for plasmid pPPS1;
  - Figure 4 shows the restriction map for plasmid pPPL1;
  - Figure 5 shows the restriction map for plasmid pATP1;
- Figure 6 shows the restriction map for plant transformation vector pVictor IV GN;
  Figure 7 shows the restriction map for plant transformation vector pVictor IV SGin Man;
  - Figure 8 shows the N terminal amino acid sequence of the rubisco activase AGP small subunit fusion enzyme, the N terminal amino acid sequence of the rubisco
- 15 activase AGP large subunit fusion enzyme and the N terminal amino acid sequence of the starch branching enzyme - AGP large subunit fusion enzyme;
  - Figure 9 shows the restriction map for plasmid pPPS4;
  - Figure 10 shows the restriction map for plasmid pBETP5;
  - Figure 11 shows the restriction map for plasmid pPPLA;
- 20 Figure 12 shows the restriction map for plasmid pPPL5;
  - Figure 13 shows the restriction map for plasmid pPPL1M;
  - Figure 14 shows the cDNA and amino acid sequences for the large sub-unit of barley AGP;
- Figure 15 shows the cDNA and amino acid sequences for the small sub-unit of barley AGP:
  - Figure 16 shows the cDNA (first 480 nucleotides from 5' end) and amino acid (first 120 amino acids from amino terminus) sequences for the starch branching enzyme; and
  - Figure 17 shows the genomic DNA sequence for the preferred potato promoter.

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In the following Examples the following amino acid codes are used:

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•		<b> </b>
	•	
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Symbol	3-letter	Meaning
Α	Ala	Alanine
В	Asp, Asn	Aspartic Asparagine
С	Cys	Cysteine
D	Asp	Aspartic
E	Glu	Glutamic
F	Phe	Phenylalanine
G	Gly	Glycine
Н	His	Histidine
I	Ile	Isoleucine
K	Lys	Lysine
L	Leu	Leucine
M	Met	Methionine
N	Asn	Asparagine
P	Pro	Proline
Q	Gln	Glutamine
R	Arg	Arginine
S	Ser	Serine
T	Thr	Threonine
v	Val	Valine
w	Тгр	Tryptophan
Х	Xxx	Unknown
Т	Tyr	Tyrosine
Z	Glu, Gln	Glutamic Glutamine
+	End	Terminator

A. Construction of plant transformation vectors containing the ADP-glucose pyrophosphorylase genes expressed from a patatin class I promoter

#### **EXAMPLE 1**

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## Plasmid pPPS1

Plasmid pPPS1 is a pBKL4 derivative containing the construction:

10 Patatin promoter - small sub-unit AGP cDNA - 35S terminator

The AGP cassette was inserted in the KpnI site of pBKL4. The additional elements introduced in the pBKL4 T-DNA by this insertion are descibed below.

Patatin promoter: The patatin promoter is a tuber specific promoter from potato (Mignery et al. 1988, Gene 62:27-44) - see SEQ. ID. No. 7.

Small subunit ADP-glucose pyrophosphorylase (bepsF2): This is a 1.8 kb cDNA fragment encoding the small subunit ADP-glucose pyrophosphorylase from barley endosperm - see SEQ ID. No. 2.

35S terminator: The CaMV 35S terminator (Odell et al. 1985, Nature 313:810-812) is fused to the bepsF2 fragment.

In more detail, a 1.8 kb BamHI cDNA fragment encoding the barley endosperm ADP glucose pyrophosphorylase small subunit (beps) was cloned in the BamHI site of plasmid pPATA 1 (Figure 1). Plasmid pPATA 1 is a derivative of plasmid pUC19 and has tuber specific patatin class I promoter ID SEQ No. 7, a polylinker cloning region, and a 35S terminator. From the resulting plasmid the 3.1 kb KpnI fragment containing the patatin promoter, the beps cDNA and the 35S terminator was isolated and inserted in the KpnI site of the plant transformation vector pBKL4 (Figure 2) to yield plasmid pPPS1 (Figure 3).

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Plasmid pBKL4 is a derivative of plasmid pBIN19 for Agrobacterium tumefaciens mediated transformation of plants and harbours a T-DNA region with a  $\beta$ -glucuronidase gene (GUS) transcribed from a 35S promoter and terminated at the nopaline synthase gene terminator, a polylinker cloning region, and a neophosphotransferase gene transcribed from a 35S promoter and terminated at the octopine synthase gene terminator.

#### EXAMPLE 2

# 10 Plasmid pPPL1

Plasmid pPPL1 is a pBKL4 derivative containing the construction:

Patatin promoter - Large subunit ADP-glucose pyrophosphorylase cDNA - 35S terminator.

The ADP-glucose pyrophophorylase cassette was inserted in the EcoRI site of pBKLA.

The additional elements introduced in the pBKL4 T-DNA by this insertion are described below.

Patatin promoter: The patatin promoter is a tuber specific promoter from potato (Mignery et al. 1988, Gene 62:27-44) - see SEQ. ID. No. 7.

Large subunit ADP-glucose pyrophosphorylase (bep110): This is a 1.9 kb cDNA fragment encoding the large subunit ADP-glucose pyrophosphorylase from barley endosperm (Villand et al. 1992, Plant Physiol 100:1617-1618) - see SEQ ID. No. 1.

35 **terminator:** The CaMV 35S terminator (Odell et al. 1985, Nature 313:810-812) is fused to the bepl 10 fragment.

In more detail, a 1.9 kb EcoRI-HindIII cDNA fragment encoding the barley endosperm ADP glucose pyrophosphorylase large subunit (bepl) was isolated, the restriction ends were filled in with klenow DNA polymerase, and the blunt ended DNA fragment was cloned in the SmaI site of plasmid pPATA1. From the resulting plasmid the 3.2 kb EcoRI fragment containing the patatin promoter, the bepl cDNA, and the 35S terminator was isolated and inserted in the EcoRI site of the plant transformation vector pBKLA to yield plasmid pPPL1 (Figure 4).

# **EXAMPLE 3**

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#### Plasmid pPPL1M

Plasmid pPPL1M (see Figure 13) is similar to pPPL1 except that the ADP-glucose pyrophosphorylase cassette:

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Patatin promoter - Large subunit ADP-glucose pyrophosphorylase - 35S terminator.

was inserted in the EcoRI site of pVictorIV SGiN Man.

#### 20 EXAMPLE 4

#### Plasmid pVictor IV SGN

pVictorIV SGN (Figure 6) is a vector for Agrobacterium mediated plant transformation, and contains the Ti right and left border sequences from the nopaline type pTiT37 plasmid (Yadav et al. 1982, Proc Natl Acad Sci 79:6322-6326) flanking the genes encoding kanamycin resistance (NPTII) and B-glucoronidase (GUS).

For replication and maintenance in *E. coli* the plasmid contains the origin of replication from the *E. coli* plasmid pUC19 (pUC19ori) Yanish-Perron et al. 1985 Gene 33:103-119), and for replication and maintenance in *Agrobacterium tumefaciens* the plasmid further contains the origin of replication from the *Pseudomonas* plasmid

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pVS1 (pVS1ori) (Itoh et al. 1984, Plasmid 11:206-220, Itoh and Haas 1985, Gene 36:27-36). For selection in *E. coli* and *Agrobacterium tumefaciens* the plasmid contains the spectinomycin/streptomycin resistance gene (spec.strep) from the transposon Tn7 encoding the enzyme 3'(9)-O- nucleotidyltransferase (Fling et al. 1985, Nucleic Acids Res 19:7095-7106). The spec/strep resistance gene is fused to the *tac* promoter for efficient expression in the bacterium.

The T-DNA segment between the right and left border harbours the following genes, which are the only genes transferred to the potato plant via the Agrobacterium tumefaciens mediated transformation.

B-glucuronidase (GUS): This segment next to the right border is the B-glucuronidase gene (GUS) from *E. coli* (Jefferson et al., 1986, Proc Natl Acad Sci 83:8447-8451) fused to the CaMV 35S promoter (35S) and 35S terminator (35St) (Odell et al. 1985, Nature 313:810-812).

Multiple cloning sites (MCS): A polylinker containing various restriction endonuclease recognition sites is inserted after the 35S terminator.

Kanamycin resistance (NPTII): The segment next to the MCS is the kanamycin (neomycin) phosphotransferase gene (NPTII) from the transposon Tn5 (Beck et al. 1982 Gene 19:327-336) fused to the CaMV 35S promoter (Odell et al. 1985, Nature 313:810-812) and the terminator of the octopine synthase gene (Caplan et al. 1983, Science 222:815-821).

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# **EXAMPLE 5**

#### Plasmid pVictor IV SGiN Man

pVictorIV SGiN Man (Figure 7) is similar to pVictorIV SGN (Bilag XI) except that the GUS gene is replaced by another GUS gene containing an intron (GUSintron) to prevent expression in bacteria.

Moreover, the kanamycin (neomycin) phosphotransferase gene (NPTII) has been replaced by the mannose-6-phosphate isomerases gene, manA, from E. coli.

- B-glucuronidase (GUSintron): This segment next to the right border is the B-glucuronidase gene (GUS) from E. coli (Jefferson et al. 1986, Proc Natl Acad Sci 83:8447-8451) furnished with an intron to prevent expression in bacteria, and fused to the CaMV 35S promoter (35S) and 35S terminator (35St) (Odell et al. 1985, Nature 313:810-812).
- 10 Mannose-6-phosphate isomerase: This segment is the mannose-6-phosphate isomerases gene, manA, from E. coli (Miles and Guest 1984, Gene 32:41-48) fused to the enhanced 35S promoter (E35S) (Kay et al. 1987, Science 236:1299-1302) and 35S terminator (35St) (Odell et al. 1985, Nature 313:810-812). The phosphomannose isomerase gene is used as a selection marker to select transgenic shoots on a media containing D-mannose as the carbon source.
  - B. Attachment of transit peptides to the ADP-Glucose pyrophosphorylase subunits

# 20 EXAMPLE 6

# Plasmid pPPS4

pPPS4 (Figure 9) is a pVictorIV derivative in which a 3 kb KpnI fragment containing the construct

Patatin promoter - spinach rubisco activase transit peptide - small subunit AGP from barley endosperm - 35S terminator

30 is inserted in the KpnI site.

In more detail, the coding region of the barley endosperm ADP glucose pyrophosphorylase small subunit (beps) cDNA was amplified by PCR using the primers:

- 5 5' CGG GAT CCA TGG ATG TAC CTT TGG CA 3' and
  - 5' CGG GAT CCT TAT TTA TTT ATA TGA CTG TTC CAC TAG 3'

which provide the PCR fragment with a BamHI and a NcoI in the 5' end and a 10 BamHI site at the 3' end. The 1.4 kb BamHI fragment containing the entire coding region of the AGP small subunit plus two additional amino acids (G and S) at the amino-terminal end was cloned in the BamHI site of pBluescript II KS to yield plasmid pBBSF. The 1.4 kb BamHI fragment was isolated from pBBSF and cloned in the BamHI site of plasmid pATP1 (Figure 5). Plasmid pATP1 has a patatin 15 promoter, a 58 amino acid rubisco activase transit peptide DNA and 35 amino acids of the mature enzyme, a BamHI site that facilitates in frame fusion of the small subunit AGP reading frame, and a 35S terminator. The 3 kb KpnI fragment including the patatin promoter, the activase transit peptide, the AGP coding region, and the 35S terminator was isolated from the resulting plasmid and cloned in the KpnI site of the plant transformation vector pVictorIV GN (Figure 6) to give plasmid 20 pPPS4 (Figure 9).

Amino terminal amino acid sequence of the rubisco activase - AGP small subunit fusion enzyme.

- 25
- 1 MATAVSTVGA ATRAPLNLNG SSAGASVPTS GFLGSSLKKH 40
- 41 TNVRFPSSSR TTSMTVKAAE NEEKNTDKWA HLAKDFSDDQ 80
- 30 81 LDIRRGKGMV DSLGSMDVPL ASKVPLPSPS KHEQCNVYSH 120

The rubisco activase sequences starts at amino acid residue 1 and ends at leucine residue at 93, while the AGP small subunit sequences begins with the methionine at 96. The rubisco activase transit peptide is cleaved at the alanine residue at 58 leaving the alanine at 59 as the N-terminal amino acid.

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The above sequence is listed later on as SEQ.I.D. No. 8.

#### **EXAMPLE 7**

#### 10 Plasmid pPPL4

pPPL4 is a pVictorIV SGiN Man derivative in which a 3.2 kb EcoRI fragment containing the construct

Patatin promoter - spinach rubisco activase transit peptide - large subunit AGP from barley endosperm - 35 S terminator

is inserted in the EcoRI site.

- 20 In more detail, the coding region of the barley endosperm ADP glucose pyrophosphorylase large subunit (bepl) cDNA was amplified by PCR using the primers
  - 5' GCG GAT CCA TAT CGA GTT CAG CGT 3'
- 25 and
  - 5' CGG GAT CCG CAC AGG TTG TCG CAG AAC 3'

which provide the PCR fragment with a BamHI and a NdeI in the 5' end and a BamHI site at the 3' end. The 1.6 kb BamHI fragment containing the entire coding region of the AGP large subunit plus two additional amino acids (I and H) at the amino-terminal end was cloned in the BamHI site of pBluescript II KS to yield plasmid pBBLF.

The 1.6 kb BamHI fragment was isolated from pBBLF and cloned in the BamHI site of plasmid pATP2. Plasmid pATP2 has a patatin promoter, a 58 amino acid rubisco activase transit peptide DNA and 35 amino acids of the mature enzyme, a BamHI site that facilitates in frame fusion of the large subunit AGP reading frame, and a 35S terminator. The 3 kb EcoRI fragment including the patatin promoter, the activase transit peptide, the AGP coding region, and the 35S terminator was isolated from the resulting plasmid and cloned in the EcoRI site of the plant transformation vector pVictor IV SGiN Man (Figure 7) to form plasmid pPPL4 (Figure 11).

- Amino terminal amino acid sequence of the rubisco activase AGP large subunit fusion enzyme.
  - 1 MATAVSTVGA ATRAPLNLNG SSAGASVPTS GFLGSSLKKH 40
  - 41 TNVRFPSSSR TTSMTVKAAE NEEKNTDKWA HLAKDFSDDQ 80
- 15 81 LDIRRGKGMV DSLGIHMQFS SVLPLEGKAC VSPVRREGSA 120

The rubisco activase sequences starts at amino acid residue 1 and ends at the leucine residue at 93, while the AGP large subunit sequences begins with the methionine at 97. The rubisco activase transit peptide is cleaved at the alanine residue at 58 leaving the alanine at 59 as the N-terminal amino acid. The above sequence is listed later on as SEQ.I.D. No. 9.

#### **EXAMPLE 8**

# 25 Plasmid pPPL5

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pPPL5 is a pVictorIV GIN MAN derivative in which a 3.4 kb EcoRI fragment containing the construct

Patatin promoter - potato starch branching enzyme amyloplast transit peptide - large subunit AGP from barley endosperm - 35S terminator

is inserted in the EcoRI site.

The coding region of the barley endosperm ADP glucose pyrophosphorylase large subunit was amplified by PCR using the primers:

- 5' CGG GAT CCG ATG CAG TTC AGC AGC GTG 3'
- 5 and

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5' CGG GAT CCG CAC AGG TTG TCG CAG AAC 3'

which provide a 1.62 kb PCR fragment with BamHI ends. The BamHI fragment containing the entire coding region of the AGP large subunit plus one additional amino acid (P) at the amino terminal end was inserted in the BamHI site of pBETP5 (Figure 10). In this way the AGP large subunit was fused to the 75 amino acid potato starch branching enzyme transit peptide plus 26 amino acids of the mature branching enzyme. The fusion enzyme is expressed from a patatin promoter and terminated at a 35S terminator. The 3.4 Kb EcoRI fragment from the resulting plasmid (pPBL1) containing the patatin promoter, the starch branching enzyme transit peptide-AGP large subunit fusion enzyme, and the 35S terminator, was inserted in the EcoRI site of the plant transformation vector pVictorIV SGiN Man yielding plasmid pPPL5 (Figure 12).

- 20 Amino terminal amino acid sequence of the starch branching enzyme AGP large subunit fusion enzyme.
  - 1 MEINFKVLSK PIRGSFPSFS PKVSSGASRN KICFPSQHST 40
  - 41 GLKFGSQERS WDISSTPKSR VRKDERMKHS SAISAVLTDD 80
- 25 81 NSTMAPLEED VKTENIGLLN LDPMOFSSVL PLEGKACVSP 120

The starch branching enzyme sequences starts at amino acid residue 1 and ends at 103, while the AGP large subunit sequence begins with the methionine at 104. The starch branching enzyme transit peptide is cleaved at the alanine residue (75) leaving the valine residue (76) as the amino terminal amino acid.

The above sequence is listed later on as SEQ.I.D. No. 10.

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# C. Production of transgenic potato plants containing the AGP-gene

#### **EXAMPLE 9**

# 5 Axenic stock cultures

Shoot cultures of Solanum tuberosum 'Bintje' and 'Dianella' are maintained on a substrate (LS) of a formula according to Linsmaier, E.U. and Skoog, F. (1965), Physiol. Plant. 18: 100-127, in addition containing 2  $\mu$ M silver thiosulphate at 25°C and 16 h light/8 h dark.

The cultures are subcultured after approximately 40 days. Leaves are cut off the shoots and cut into nodal segments (approximately 0.8 cm) each containing one node.

# 15 Inoculation of potato tissues

Shoots from approximately 40 days old shoot cultures (height approximately 5-6 cms) were cut into internodal segments (approximately 0.8 cm). The segments are placed into liquid LS-substrate containing the transformed Agrobacterium tumefaciens containing the binary vector of interest. The Agrobacterium are grown overnight in YMB-substrate (dipotassiumhydrogen phosphate, trihydrate (0.66 g/l); magnesium sulphate, heptahydrate (0.20 g/l); sodium chloride (0.10 g/l); mannitol (10.0 g/l); and yeast extract (0.40 g/l)) containing appropriate antibiotics (corresponding to the resistance gene of the Agrobacterium strain) to an optical density at 660 nm (OD-660) of approximately 0.8, centrifuged and resuspended in the LS-substrate to an OD-660 of 0.5.

The segments are left in the suspension of <u>Agrobacterium</u> for 30 minutes and then the excess of bacteria are removed by blotting the segments on sterile filter paper.

#### Co-cultivation

The shoot segments are co-cultured with bacteria for 48 hours directly on LS-substrate containing agar (8.0 g/l), 2,4-dichlorophenoxyacetic acid (2.0 mg/l) and trans-zeatin (0.5 mg/l). The substrate and also the explants are covered with sterile filter papers, and the petri dishes are placed at 25°C and 16 h light/8 dark.

## "Washing" procedure

After the 48 h on the co-cultivation substrate the segments are transferred to containers containing liquid LS-substrate containing 800 mg/l carbenicillin. The containers are gently shaken and by this procedure the major part of the Agrobacterium are washed off the segments and/or killed.

#### 15 Selection

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After the washing procedure the segments are transferred to plates containing the LS-substrate, agar (8 g/l), trans-zeatin (1-5 mg/l), gibberellic acid (0.1 mg/l), carbenicillin (800 mg/l), and kanamycin sulphate (50-100 mg/l) or phosphinotricin (1-5 mg/l) or mannose (5 g/l) depending on the vector construction used.

The segments are sub-cultured to fresh substrate each 3-4 weeks.

In 3 to 4 weeks, shoots develop from the segments and the formation of new shoots continues for 3-4 months.

#### Rooting of regenerated shoots

The regenerated shoots are transferred to rooting substrate composed of LS-substrate, agar (8 g/l) and carbenicillin (800 mg/l).

The transgenic genotype of the regenerated shoot are verified by testing the rooting ability on the above mentioned substrates containing kanamycin sulphate (200 mg/l), by performing NPTII assays (Radke, S. E. et al, Theor. Appl. Genet. (1988), 75: 685-694) or by performing a GUS assay on the co-introduced  $\beta$ -glucuronidase gene according to Hodal, L. et al. Pl. Sci. (1992), 87: 115-122 or by assaying the for the expression of the barley AGP mRNA or AGP enzyme activity as described elsewhere. Plants which are not positive in any of these assays are discarded or used as controls.

#### Transfer to soil

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The newly rooted plants (height approx. 2-3 cms) are transplanted from rooting substrate to soil and placed in a growth chamber (21°C, 16 hour light 200-400uE/m²/sec).

When the plants are well established they are transferred to the greenhouse, where they are grown until tubers have developed and the upper part of the plants are senescing.

#### **Harvesting**

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The potatoes were harvested after about 3 months.

#### AGP Assay

25 Tubers from the harvested potato plants were stored at 4°C. AGPase was extracted by homogenization of 10-20g of thinly sliced potato tubers in 20 ml of buffer containing 25 mM Hepes (pH 7.4) mM mercaptoethanol and 1 mM DTT. Homogenization was performed at 0-4°C using 30 ml Waring blender at full speed for 15 seconds. Aliquots of crude extract were then immediately centrifuged at maximal speed for 1 min using bench Eppendorf microcentrifuge and then assayed for AGPase activity. Assays were carried out immediately after centrifugation to make sure that the enzyme will not be inactivated during storage.

Assays were carried out in the pyrophosphorolysis direction monitoring glucose-1-P formation at 340 nm (21°C), using LKB spectrophotometer (Ultrospec II). Assay mixtures (1 ml) contained: 100 mM Mops (pH 7.4), 0.6 mM NAD, 7mM MgCl<sub>2</sub>, 1 mM ADP-glucose, 1 mM inorganic pyrophosphate, 10 uM glucose-1,6-biphosphate, 2 units each of glucose-6-P dehydrogenase and phosphoglucomutase. In some instances, 2 mM 3-phosphoglyceric acid (PGA) was added to assays. Assays were run (-PGA) for ca. 5 min, and then PGA was added and assays were monitored for another 5-10 min. Rates were usually linear during the time-course of assays. One unit of AGPase activity corresponds to the amount of enzyme producing 1 umole of NADH under assay conditions.

#### Starch Analysis

The starch contents of potato tubers was determined according to a method which was designed and proposed by the Dutch-German working group "Standardization" and published in "Methods of assessment for potatoes and potato products". The method was developed for use with a sample size of 5000 g but we scaled the method down for use with potato tubers from a single plant, usually between 20 and 200 g.

All potatoes from a plant are washed and dried with a cloth before weighing (a grams) on an electronic balance. Later, the tubers are weighed again, but this time on a balance with two metal baskets of which one is immersed in a water basin. The potato tubers are placed in the bucket in water, and their weight (b grams) in water is determined.

25

The under-water weight of a sample is calculated at 5000 b/a grams. From tables showing the relation between under-water weight, dry matter and starch content, the two latter figures can be determined.

This procedure is described in more detail by W.A. Gould in Chipping Potato Handbook, ed. Gould, W.A. The Snack Food Association, Vermont, 1989, pp 18-22, in an article entitled "Specific gravity, its measurement and use.

#### **RESULTS**

AGPase levels and starch levels were increased with the constructs of the present invention, particularly in the absence of PGA, especially with the constructs coding for the large sub-unit and in particular the constructs coding for a transit peptide.

In this regard, some results are shown in Table I (below) for transformed potatoes comprising constructs derived from plasmids pPPS4 and pPPL4.

TABLE I
STARCH CONTENT IN TRANSFORMED POTATOES (TRANS)
COMPARED TO CONTROL NATIVE POTATOES
GROWN UNDER THE SAME CONDITIONS

٠.		•	•
15	Sample	% Overall	%Starch Content vis-a-vis
		Starch Content	Control
	Control	16	100
	Trans 1	19	119
20	Trans 2	23	144
	Trans 3	26	163
	Trans 4	19	119
	Trans 5	26	163
	Trans 6	24	150
25	Trans 7	21	131
	Trans 8	17	106
	Trans 9	21	131

The above results clearly show that the average starch level in the transgenic potatoes comprising constructs derived from plasmids pPPS4 and pPPL4 are increased to about 136 % of that found in native potatoes.

Furthermore, the average starch level for the upper two quartiles for the transgenic potatoes comprising constructs derived from plasmids pPPS4 and pPPL4 is about 155 % of that found in native potatoes.

- 5 Each of these findings is significant.
- 10 Other modifications of the present invention will be apparent to those skilled in the art without departing from the scope of the invention.

### SEQUENCE LISTING

### (1) GENERAL INFORMATION

5 NAME OF APPLICANTS: DANISCO A/S

**BUSINESS ADDRESS:** 

Langebrogade 1

DK-1001 Copenhagen K

Denmark

10 TITLE OF INVENTION: TRANSGENIC ORGANISM

(2a) INFORMATION FOR SEQUENCE I.D. 1

SEQUENCE TYPE:

**NUCLEIC ACID** 

15 MOLECULE TYPE:

DNA

ORIGINAL SOURCE:

BARLEY

SEQUENCE LENGTH: STRANDEDNESS:

2037

TOPOLOGY:

DOUBLE LINEAR

20 SEQUENCE:

Nucleotide sequence of a cDNA encoding the large

subunit of ADP-glucose pyrophosphorylase from barley

seed endosperm (bepl10)

#### 25 SEQ. ID. No. 1

35

40

45

1 ACGACCACCT CCGAACTCAA CGCCTCCACG GACCATCTCT

41 CTCCTCTCCC CTCCCCTCAC CACCACCACC ACCACCACCC

81 CTTCTCCCTC CCTGCATTTG ATTCGTTCAT ATTCATCCGT

30 121 CGCTTGCCCG GTCGCCACCC CGTCGATCCC TCACCCCGCC

161 GTCCCCGGCA GTTGCAGGTG GACTGCTAAT GTCATCGATG

201 CAGTTCAGCA GCGTGCTGCC CCTGGAGGCC AAGGCGTGCG

241 TITCCCCAGT CAGGAGAGAG GGATCGGCCT GCGAGCGCCT

281 CAAGATCGGG GACAGCAGCA GCATCAGGCA CGAGAGAGCG

321 TCCAGGAGGA TGTGCAACGG CGGCGCAGGG GCCCCGCCGC

361 CACCGGTGCG CAGTGCGTGC TCACCTCCGA CGCCAGCCCG

401 GCCGACACCC TTGTTCTCCG GACGTCCTTC CGGAGGAATT ACGCCGATCC GAACGAGGTC GCGGCCGTCG GTCGCGGCCG

TCATACTCGG CGGCGCACC GGGACTCAGC TCTTCCCGCT CACAAGCACA AGGGCCACAC CTGCTGTTCC TATTGGAGGA

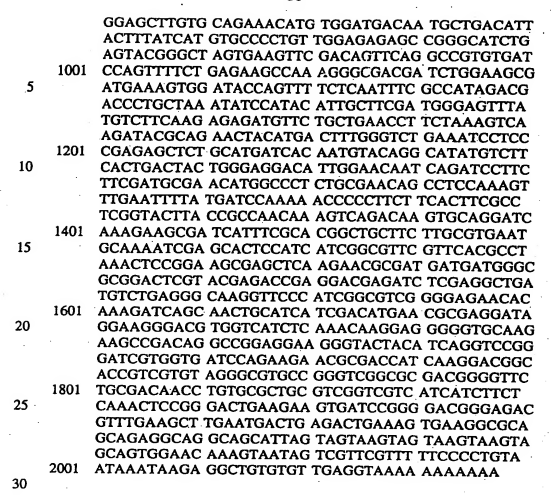
TGTTACAGGC TCATCGATAT TCCCATGAGC AACTGCTTCA

601 ACAGTGGCAT CAACAAGATA TTCGTCATGA CCCAGTTCAA
CTCGGCATCT CTCAATCGCC ACATTCACCG CACCTACCTC

GGCGGGGAA TCAATTICAC TGATGGATCT GTTGAGGTAT TGGCCGCGAC ACAAATGCCT GGGGAGGCTG CTGGATGGTT

CCGCGGAACA GCGGATGCCG TCAGAAAATT TATCTGGGTG

801 CTTGAGGACT ACTATAAGCA TAAATCCATA GAGCACATTT
TGATCTTGTC GGGCGATCAG CTTTATCGCA TGGATTACAT



10

### (2b) INFORMATION FOR SEQUENCE I.D. 2

SEQUENCE TYPE:

**NUCLEIC ACID** 

MOLECULE TYPE:

DNA

**ORIGINAL SOURCE:** SEQUENCE LENGTH:

BARLEY 1822

STRANDEDNESS:

DOUBLE

TOPOLOGY:

LINEAR

SEQUENCE:

Nucleotide sequence of a cDNA encoding the small subunit of ADP-glucose pyrophosphorylase from barley

seed endosperm (beps)

COMMENT:

The "." at 1569 denotes a purine.

SEQ. ID. No. 2

15

1 AAAAGTGAAC TCACACATCA CTCAATATCT ATATCCTTCC ATTITATATC CCTCGGTGAT GGATGTACCT TTGGCATCTA AAGTTCCCTT GCCCTCCCCT TCCAAGCATG AACAATGCAA CGTTTATAGT CATAAGAGCT CATCGAAGCA TGCAGATCTC 20 AATCCCCATG CTATTGATAG TGTTCTCGGT ATCATTCTTG GAGGTGGTGC AGGGACTAGA TTGTATCCCC TGACGAAGAA 201 GCGTGCAAAG CCTGCAGTGC CATTGGGTGC CAACTACAGG CITATTGATA TTCCTGTCAG TAATTGTCTG AACAGCAACA TATCAAAGAT CTATGTGCTT ACACAGTTCA ACTCAGCTTC TCTTAATCGT CATCTCTCAC GAGCCTATGG GAGCAACATT 25 GGAGGTTACA AGAATGAAGG ATTTGTTGAA GTCCTTGCTG 401 CACAGCAGAG CCCAGATAAC CCTGACTGGT TCCAGGGTAC TGCAGATGCT GTAAGGCAGT ACTTGTGGCT ATTCGAGGAG CATAATGTTA TGGAGTATCT AATTCTTGCT GGAGATCACC 30 TGTACCGAAT GGACTATGAA AAGTTTATTC AGGCACACAG 601 AGAAACGGAT GCTGATATTA CTGTTGCTGC CTTGCCCATG GATGAGGAAC GTGCAACTGC ATTTGGCCTT ATGAAAATCG ATGAAGAAGG GAGGATAATT GAATTCGCAG AGAAACCAAA AGGAGAACAG TTGAAAGCTA TGATGGTTGA TACGACCATA 35 CTTGGCCTTG AAGATGCGAG GGCAAAGGAA ATGCCTTATA TTGCTAGCAT GGGTATCTAT GTTATTAGCA AACATGTGAT 801 GCTTCAGCTT CTCCGTGAGC AATTTCCTGG AGCTAATGAC TTCGGAAGTG AAGTTATCCC TGGTGCAACT AGCACTGGCA TGAGGGTACA AGCATACCTA TACGACGGTT ACTGGGAAGA 40 TATTGGTACA ATTGAGGCAT TCTATAATGC AAATTTGGGA 1001 ATTACCAAAA AACCAATACC TGATTTCAGT TTCTATGACC GTTCTGCTCC CATTTACACA CAACCTCGAC ACTTGCCTCC TTCAAAGGTT CTTGATGCTG ATGTGACAGA CAGTGTAATT GGTGAAGGAT GTGTTATTAA AAACTGCAAG ATACACCATT 45 CAGTAGTTGG ACTCCGTTCC TGCATATCTG AAGGTGCAAT AATAGAGGAC ACGTTGCTAA TGGGTGCGGA CTACTATGAG 1201

ACTGAAGCTG ATAAGAAACT CCTTGCTGAA AAAGGTGGCA TTCCCATTGG TATTGGAAAG AATTCACACA TCAAAAGAGC

		AATCATTGAC AAGAATGCTC GTATTGGAGA TAACGTGATG
		ATAATCAATG TTGACAATGT TCAAGAAGCG GCGAGGGAGA
	1401	CAGATGGATA CTTCATCAAA AGTGGCATCG TAACTGTGAT
		CAAGGATGCT TTACTCCCTA GTGGAACAGT CATATGAAGC
5	•	AGATGTGAAA TGTATGCCAA AAGACAGGGC TACTTGCGTC
		AGTCTGGAAT CAACCAACAA GGCCGCGAAG GAGATCATAA
		AATAAAAA.G GAGTGCCATG CGAGTCACTT CTACACCCTT
	1601	TTCCCCCCTT GATGTATTAG GAACTGTGAT GTACAAGCAA
		CTGTGATGCA CTTACGCGAA GTGCCCCTGG ATTCAGCTTT
10		CTCTTTGCTT GTAACTGGTT TCCAGCAGAC CATGCTATTT
		GTTGTATGGT TCGTGCAAAA CCTTGCGATG CTTTATATAT
		GCTTTATATA TAAACAAGAT GAATCCCCGC GCGTTGCTGC
	2001	GGCACAAAA AAAAAAAAA AA

## (2c) INFORMATION FOR SEQUENCE I.D. 3

SEQUENCE TYPE:

**ENZYME** 

MOLECULE TYPE:

AMINO ACID

ORIGINAL SOURCE:

BARLEY

SEQUENCE LENGTH:

528

TOPOLOGY:

LINEAR

SEQUENCE:

Derived amino acid sequence of a cDNA encoding the large subunit of ADP-glucose pyrophosphorylase from

barley seed endosperm (bepl10)

10

SEQ. ID. No. 3

	1	MSSMQFSSVL	PLEGKACVSP
15		VRREGSACER	LKIGDSSSIR
		HERASRRMCN	GGAGAPPPV
		RSACSPPTPA	RPTPLFSGRP
		SGGITPIRTR	SRPSVAAVIL
•	101		LTSTRATPAV
20		PIGGCYRLID	IPMSNCFNSG
		INKIFVMTQF	NSASLNRHIH
٠.		RTYLGGGINF	
		TQMPGEAAGW	TDGSVEVLAA
ė.	201	FIWVLEDYYK	FRGTADAVRK
25		SGDQLYRMDY	HKSIEHILIL
	•	NADITLSCAP	MELVQKHVDD
		LVKFDSSGRV	VGESRASEYG
		DLEAMKVDTS	IQFSEKPKGD
	301	KYPYIASMGV	FLNFAIDDPA
30	,	LLKSRYAELH	Y V F K R D V L L N D F G S E I L P R A
		LHDHNVQAYV	
		IRSFFDANMA	FTDYWEDIGT
		YDPKTPFFTS	LCEQPPKFEF
	401		PRYLPPTKSD
35	401	EHSIIGVRSR	HGCFLRECKI
-		MMMGADSYET	LNSGSELKNA
		GKVPIGVGEN	EDEISRLMSE
		NARIGRDVVI	TKISNCIIDM
	<b>501</b>	RPEEGYYIRS	SNKEGVQEAD
40	501	IKDGTVV*	GIVVIQKNAT

### (2d) INFORMATION FOR SEQUENCE I.D. 4

SEQUENCE TYPE:

**ENZYME** 

MOLECULE TYPE:

AMINO ACID

5 ORIGINAL SOURCE:

**BARLEY** 

SEQUENCE LENGTH:

472

TOPOLOGY:

LINEAR

**SEQUENCE:** 

Derived amino acid sequence of a cDNA encoding the

small subunit of ADP-glucose pyrophosphorylase from

barley seed endosperm (beps)

SEQ. ID. No. 4

15	1	DVPLASKVPL	PSPSKHEOCN
•		VYSHKSSSKH	ADLNPHAIDS
		VLGIILGGGA	
		RAKPAVPLGA	
		NCLNSNISKI	
20	101		YVLTQFNSAS
20	101	LNRHLSRAYG	SNIGGYKNEG
•		FVEVLAAQQS	PDNPDWFQGT
		ADAVRQYLWL	FEEHNVMEYL
		ILAGDHLYRM	DYEKFIQAHR
		ETDADITVAA	LPMDEERATA
25	201	FGLMKIDEEG	RIIEFAEKPK
		GEQLKAMMVD	TTILGLEDAR
		AKEMPYIASM	GIYVISKHVM
	•	LQLLREQFPG	
		GATSTGMRVQ	ANDFGSEVXP
30	301		AYLYDGYWED
50	301	IGTIEAFYNA	NLGITKKPIP
		DFSFYDRSAP	IYTQPRHLPP
		SKVLDADVTD	SVIGEGCVIK
		NCKIHHSVVG	LRSCISEGAI
		IEDTLLMGAD	YYETEADKKL
35	401	LAEKGGIPIG	IGKNSHIKRA
		IIDKNARIGD	NVMIINVDNV
		QEAARETDGY	FIKSGIVTVI
		KDALLPSGTV	I *
		A D A D L P 3 O I V	1 -

40

### (2e) INFORMATION FOR SEQUENCE I.D. 5

SEQUENCE TYPE:

**NUCLEIC ACID** 

MOLECULE TYPE:

DNA **POTATO** 

**ORIGINAL SOURCE: SEQUENCE LENGTH:** 

480

STRANDEDNESS: TOPOLOGY:

**DOUBLE** 

LINEAR

SEQUENCE:

Starch branching enzyme cDNA

10

5

(First 480 nucleotides from 5' end)

SEQ. ID. No. 5

CCCGTCTGTA AGCATCATTA GTGATGTTGT 1 15 31 TCCAGCTGAA TGGGATGATT CAGATGCAAA 61 CGTCTGGGGT GAGAACATAC AAGAAGGCAG 20 91 CAGCTGAAGC AAAGTACCAT AATTTAATCA 121 ATGGAAATTA ATTTCAAAGT TTTATCAAAA 151 CCCATTCGAG GATCTTTTCC ATCTTTCTCA 25 CCTAAAGTTT CTTCAGGGGC TTCTAGAAAT AAGATATGTT TTCCTTCTCA ACATAGTACT 30 GGACTGAAGT TTGGATCTCA GGAACGGTCT TGGGATATTT CTTCCACCCC AAAATCAAGA 301 GTTAGAAAG ATGAAAGGAT GAAGCACAGT 35 TCAGCTATTT CCGCTGTTTT GACCGATGAC AATTCGACAA TGGCACCCCT AGAGGAAGAT 40 GTCAAGACTG AAAATATTGG CCTCCTAAAT TTGGATCCAA CTTTGGAACC TTATCTAGAT 451 CACTTCAGAC ACAGAATGAA GAGATATGTG 45

### (2f) INFORMATION FOR SEQUENCE I.D. 6

SEQUENCE TYPE:

PEPTIDE

MOLECULE TYPE:

AMINO ACID

ORIGINAL SOURCE: SEQUENCE LENGTH:

**POTATO** 

ACCOUNTED TO

120

TOPOLOGY:

LINEAR

SEQUENCE:

Starch branching enzyme amino acid

(First 120 amino acids from amino terminus)

10

5

SEQ. ID. No. 6

- MET Glu Ile Asn Phe Lys Val Leu Ser Lys 15 11 Pro Ile Arg Gly Ser Phe Pro Ser Phe Ser 21 Pro Lys Val Ser Ser Gly Ala Ser Arg Asn 20 31 Lys Ile Cys Phe Pro Ser Gln His Ser Thr 41 Gly Leu Lys Phe Gly Ser Gln Glu Arg Ser 51 Trp Asp Ile Ser Ser Thr Pro Lys Ser Arg 25 61 Val Arg Lys Asp Glu Arg MET Lys His Ser 71 Ser Ala Ile S er Ala Val Leu Thr Asp Asp 30 81 Asn Ser Thr MET Ala Pro Leu Glu Glu Asp 91 Val Lys Thr Glu Asn Ile Gly Leu Leu Asn 101 Leu Asp Pro Thr Leu Glu Pro Tyr Leu Asp 35 111 His Phe Arg His Arg MET Lys Arg Tyr Val
- In the event that this sequence contains an error, see the corresponding sequence in the accompanying figures.

### (2g) INFORMATION FOR SEQUENCE I.D. 7

**SEQUENCE TYPE:** 

**NUCLEIC ACID** 

MOLECULE TYPE:

DNA

ORIGINAL SOURCE: SEQUENCE LENGTH:

POTATO

STRANDEDNESS:

1047 DOUBLE

STRANDEDNESS: TOPOLOGY:

LINEAR

SEQUENCE:

Tuber specific class 1 promoter

10 SEQ. ID. No. 7

1 TTGTTAGTTA ATGCGTATTA GTTTTAGCGA CGAAGCACTA AATCGTCTTT GTATACTTTC AGTGACACAT GTTTAGTGAC 15 GACTGATTGA CGAAATTTTT TTCGTCTCAC AAAATTTTTA GTGACGAAAC ATGATTTATA GATGACGAAA TTATTTGTCC CTCATAATCT AATTTGTTGT AGTGATCATT ACTCCTTTGT TTGTTTTATT TGTCATGTTA GTTCATTAAA AAAAAATCT 201 CTCTTCTTAT CAATTCTGAC GTGTTTAATA TCATAAGATT AAAAAATATT TTAATATATC TITAATTTAA AGCCACAAAA 20 TTTAAATTTC TTCGTTAACA TAATTTGTCA AATCAGGCTC AAAGATCGTT TTTCATATCG GAATCAGGAT TTTATTTATT CTTTTAAAAA TAAAGAGGTG GTGAGCTAAA CAATTTCAAA 401 TCTCATCACA CATATGGGGT CAGCCACAAA AATAAAGAAC 25 GGTTGGAACG GATCTATTAT ATAATACTAA TAAAGAATAG AAAAAGGAAA GTGAGTGAGG TGCGAGGGAG AGAATCTGTT TAATATGCAG AGTCGATCAT GTGTCAGTTT TATCGATATG ACTCTGATTT CAACTGAGTT TAAGCAATTC TGATAAGGCG 601 AGGAAAATCA CAGTGCTGAA ATCTAGAAAA ATCTCATACA 30 GTGAGATAAA TCTCAACAAA AACGTTGAGT CCATAGAGGG GGTGTATGTG ACACCCAACC TCAGCAAAAG AAAACCTCCC CTCAAGAAGG ACATTTGCGG TGCTAAACAA TTTCAAGTCT CATCACACAT ATATATTATA TAATACTAAT AAAGAATAGA 801 AAAAGGAAAG GTAAACATCA CTAATGACAG TTGCGGTGCA AAGTGAGTGA GATAATAAAC ATCAGTAATA GACATCACTA 35 ACTITIATIG GITATGICIT TCTCAAAATA AAATITCTCA ACTTGTTTAC GTGCCTATAT ATACCATGCT TGTTATATGC TCAAAGCACC AACAAAATTT AAAAACACTT TGAACATTTG 1001 **CCCCGGG** 40

45

### (2h) INFORMATION FOR SEQUENCE I.D. 8

SEQUENCE TYPE:

**ENZYME CONSTRUCT** 

MOLECULE TYPE:

AMINO ACID

5 ORIGINAL SOURCE:

SPINACH and BARLEY

**SEQUENCE LENGTH:** 

120

TOPOLOGY:

LINEAR

SEQUENCE:

Amino terminal amino acid sequence of the rubisco

activase - AGP small subunit fusion enzyme.

10

### SEQ.ID.NO.8

1 MATAVSTVGA ATRAPLNLNG SSAGASVPTS GFLGSSLKKH 40

15 41 TNVRFPSSSR TTSMTVKAAE NEEKNTDKWA HLAKDFSDDQ 80

81 LDIRRGKGMV DSLGSMDVPL ASKVPLPSPS KHEQCNVYSH 120

# (2i) INFORMATION FOR SEQUENCE I.D. 9

**SEQUENCE TYPE:** 

**ENZYME CONSTRUCT** 

MOLECULE TYPE:

AMINO ACID

5 ORIGINAL SOURCE: SPINACH and BARLEY

SEQUENCE LENGTH:

120

TOPOLOGY:

LINEAR

SEQUENCE:

Amino terminal amino acid sequence of the rubisco

activase - AGP large subunit fusion enzyme.

10

#### SEQ.ID.NO.9

MATAVSTVGA ATRAPLNLNG SSAGASVPTS GFLGSSLKKH 40 TNVRFPSSSR TTSMTVKAAE NEEKNTDKWA HLAKDFSDDQ 80 15 41

LDIRRGKGMV DSLGIHMQFS SVLPLEGKAC VSPVRREGSA 120 81

WO 94/24292 PCT/EP94/01082

- 48 -

### (2j) INFORMATION FOR SEQUENCE I.D. 10

**SEQUENCE TYPE:** 

**ENZYME CONSTRUCT** 

MOLECULE TYPE:

**AMINO ACID** 

5 ORIGINAL SOURCE:

SPINACH and BARLEY

SEQUENCE LENGTH:

120

TOPOLOGY:

LINEAR

SEQUENCE:

Amino terminal amino acid sequence of the starch

branching enzyme - AGP large subunit fusion enzyme.

10

#### SEQ.ID.NO.10

1 MEINFKVLSK PIRGSFPSFS PKVSSGASRN KICFPSQHST 40

15 41 GLKFGSQERS WDISSTPKSR VRKDERMKHS SAISAVLTDD 80

81 NSTMAPLEED VKTENIGLLN LDPMQFSSVL PLEGKACVSP 120

# INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description		
on page 19 , line 6		
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industria	l and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country	)	
23. St. Machar Drive Aberdeen Scotland United Kingdom AB2 1RY		
Date of deposit	Accession Number	
29 March 1994	NCIMB 40618	
C. ADDITIONAL INDICATIONS (leave blank if not applicable	le) This information is continued on an additional sheet	
other designated state having equivalent legislation, a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample. (Rule 28(4) EPC).  D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)		
	, and a second s	
E. SEPARATE FURNISHING OF INDICATIONS (leave	blank if not applicable)	
	Bureau later (specify the general nature of the indications e.g., "Accession	
For receiving Office use only	For International Bureau use only	
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A. The indications made below relate to the microorganism referred to in the description on page, line		
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industria	l and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country	y)	
23. St. Machar Drive	•	
Aberdeen		
Scotland United Kingdom		
AB2 1RY		
Date of deposit	Accession Number	
29 March 1994	NCIMB 40619	
C. ADDITIONAL INDICATIONS (leave blank if not applicate	ble) This information is continued on an additional sheet	
grant of the European patent or until t refused or withdrawn or is deemed to be sample to an expert nominated by the pe EPC).	til the publication of the mention of the he date on which the application has been withdrawn, only by the issue of such a rson requesting the sample. (Rule 28(4)	
	(,, )	
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E. SEPARATE FURNISHING OF INDICATIONS (leave	.11.17	
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A The indications and below the state of		
A. The indications made below relate to the microorganism r on page	eferred to in the description	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industria	l and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country	y) (v	
23. St. Machar Drive		
Aberdeen		
Scotland	·	
United Kingdom AB2 1RY		
Date of deposit	Accession Number	
29 March 1994	NCIMB 40620	
C. ADDITIONAL INDICATIONS (leave blank if not applical	ble) This information is continued on an additional sheet	
In respect of those designations in which a European patent is sought, and any other designated state having equivalent legislation, a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample. (Rule 28(4) EPC).		
D. DESIGNATED STATES FOR WHICH INDICATION	ONS ARE MADE (if the indications are not for all designated States)	
	THE HANDE (i) the thetetations are not for all aesignated states)	
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E. SEPARATE FURNISHING OF INDICATIONS (leave	: blank if not applicable)	
The indications listed below will be submitted to the International	Bureau later (specify the general nature of the indications e.g., "Accession	
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A 50 Section and Laboratory		
A. The indications made below relate to the microorganism referred to in the description on page		
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industria	l and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country	)	
23. St. Machar Drive		
Aberdeen		
Scotland		
United Kingdom AB2 1RY	and the second s	
Date of deposit	Accession Number	
29 March 1994	NCIMB 40621	
C. ADDITIONAL INDICATIONS (leave blank if not applicable	This information is continued on an additional sheet	
other designated state having equivalent microorganism will be made available ungrant of the European patent or until the refused or withdrawn or is deemed to be sample to an expert nominated by the perEPC).	til the publication of the mention of the he date on which the application has been withdrawn, only by the issue of such a	
D. DESIGNATED STATES FOR WHICH INDICATIO	NS ARE MADE (if the indications are not for all designated States)	
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	·	
E. SEPARATE FURNISHING OF INDICATIONS (leave	blank if not applicable)	
The indications listed below will be submitted to the International Number of Deposit")	Bureau later (specify the general nature of the indications e.g., "Accession	
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Zi application	This sheet was received by the International Bureau on:	
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A. The indications made below relate to the microorganism re on page, line	terred to in the description	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industria	l and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country	)	
23. St. Machar Drive	•	
Aberdeen		
· Scotland United Kingdom		
AB2 1RY		
Date of deposit	Accession Number	
29 March 1994	NCIMB 40622	
C. ADDITIONAL INDICATIONS (leave blank if not applicab	le) This information is continued on an additional sheet	
microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample. (Rule 28(4) EPC).  D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)		
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PCT/EP94/01082

#### INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page		
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industrial	and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country,		
23. St. Machar Drive		
Aberdeen Scotland	·	
United Kingdom		
AB2 1RY		
Date of deposit	Accession Number	
31 March 1994	NCIMB 40623	
C. ADDITIONAL INDICATIONS (leave blank if not applicable)	le) This information is continued on an additional sheet	
other designated state having equivalent legislation, a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample. (Rule 28(4) EPC).		
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# INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism re on page, line	ferred to in the description
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution	
The National Collections of Industrial	l and Marine Bacteria Limited (NCIMB)
Address of depositary institution (including postal code and country,	)
23. St. Machar Drive Aberdeen Scotland United Kingdom AB2 1RY	
Date of deposit	Accession Number
31 March 1994	NCIMB 40624
C. ADDITIONAL INDICATIONS (leave blank if not applicable	This information is continued on an additional sheet
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R.L.R. PETHER	
m PCT/RO/134 (July 1992)	

### INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

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A. The indications made below relate to the microorganism referred to in the description		
on page, line	25	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet	
Name of depositary institution		
The National Collections of Industria	l and Marine Bacteria Limited (NCIMB)	
Address of depositary institution (including postal code and country	y)	
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Aberdeen Scotland		
United Kingdom		
AB2 1RY	•	
Date of deposit	Accession Number	
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C. ADDITIONAL INDICATIONS (leave blank if not applical	ble) This information is continued on an additional sheet	
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D. DESIGNATED STATES FOR WHICH INDICATION	ONS ARE MADE (if the indications are not for all designated States)	
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TO POTTO (174 (Ind., 1992)		

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#### **CLAIMS**

1. A transgenic starch producing organism comprising a nucleotide sequence coding for an exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme, wherein the nucleotide sequence is capable of being expressed in the organism;

characterised in that the activity of the enzyme or sub-unit thereof is substantially independent of any level of *in vivo* 3-phospho-glycerate and/or any *in vivo* level of inorganic phosphate; and

further characterised in that the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

- 2. An organism according to claim 1 wherein the AGP enzyme is a heteromer, preferably a heterotetramer, more preferably a heteromer containing two large subunits and two small sub-units.
- 3. An organism according to claim 2 wherein the AGP enzyme is isolatable from Hordeum, preferably wherein the enzyme is barley (Hordeum vulgare) endosperm AGP or a sub-unit thereof, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units.
- 4. An organism according to claim 3 wherein the nucleotide sequence is a DNA sequence.
- An organism according to claim 4 wherein the DNA sequence encodes the large sub-unit of the barley endosperm AGP or a variant thereof having non-critical
   amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequence defining the large sub-unit of the barley endosperm AGP.

- 6. An organism according to claim 4 or claim 5 wherein the DNA sequence is the sequence identified herein as SEQ ID No 1, including non-critical allelic variations of that sequence.
- 5 7. An organism according to claim 4 wherein the DNA sequence encodes the small sub-unit of the barley endosperm AGP, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequence defining the small sub-unit of the barley endosperm AGP.
- 10 8. An organism according to claim 4 or claim 7 wherein the DNA sequence is the sequence identified herein as SEQ ID No 2, including non-critical allelic variations of that sequence.
- 9. An organism according to claim 4 comprising a DNA sequence according to claim 5 or claim 6 and a DNA sequence according to claim 7 or claim 8, preferably wherein each DNA sequence additionally codes for a different marker.
  - 10. An organism according to any one of the preceding claims wherein the expressed AGP enzyme or sub-unit thereof comprises the amino acid sequence set out in SEQ ID No. 3, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.
- 11. An organism according to any one of claims 1 to 9 wherein the expressed AGP enzyme or sub-unit thereof comprises the amino acid sequence set out in SEQ
  25 ID No. 4, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.
  - 12. An organism according to claim 1 wherein the expressed AGP comprises both a large sub-unit having the amino acid sequence set out in SEQ ID No. 3 or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence, and a small sub-unit having the amino acid sequence set out in SEQ ID No. 4 or a variant thereof having non-critical amino acid

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substitution(s) or deletion(s) at one or more locations in that sequence.

- 13. An organism according to any one of the preceding claims wherein the nucleotide sequence additionally codes for a transit peptide.
- 14. An organism according to claim 13 wherein the transit peptide is Rubisco Activase transit peptide or Starch Branching enzyme transit peptide.
- 15. An organism according to claim 14 wherein the transit peptide is coded for by
   10 a DNA sequence comprising the sequence identified as SEQ. I.D. No. 5, including non-critical allelic variations of that sequence.
  - 16. An organism according to claim 14 wherein the transit peptide has an amino acid sequence comprising the sequence identified as Seq.I.D.No. 6, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.
  - 17. An organism according to claim 4 wherein the nucleotide sequence is operatively connected to a promoter which expresses the nucleotide sequence wherein the promoter is cell, tissue or organ specific.
    - 18. An organism according to claim 17 wherein the promoter has the sequence identified as SEQ.I.D.No.7, or a variant thereof having non-critical nucleotide substitution(s) or deletion(s) at one or more locations in that sequence.
    - 19. A transgenic starch producing organism comprising exogenous ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof which retains the enzymatic activity of the AGP enzyme, wherein the activity of the enzyme or sub-unit thereof is substantially independent of any level of *in vivo* 3-phospho-glycerate and/or any *in vivo* level of inorganic phosphate and wherein the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

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- 20. An organism according to claim 19 wherein the AGP enzyme or sub-unit thereof comprises the amino acid sequence set out in SEQ ID No. 3, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence and/or the amino acid sequence set out in SEQ ID No. 4, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.
- 21. An organism according to any one of the preceding claims wherein the organism is a transgenic plant.

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- 22. An organism according to claim 20 wherein the transgenic plant is a potato plant.
- 23. A potato tuber containing an enhanced starch content, when obtained from a transgenic potato plant according to claim 22.
  - 24. A method of increasing the rate and/or yield of starch production in an organism, especially a plant or a plant cell, which method comprises introducing into an organism a nucleotide sequence as defined in claim 1 or any claim dependent thereon to form a transgenic organism as defined in any one of the preceding claims and expressing the nucleotide sequence.
- 25. A method of increasing the rate and/or yield of starch production in an organism, especially a plant or a plant cell, which method comprises introducing into
   25 or forming in an organism a ADP glucose pyrophosphorylase (AGP) enzyme or a sub-unit thereof as defined in claim 19 or any claim dependent thereon.
  - 26. A cDNA sequence identified herein as SEQ ID No. 2, including non-critical allelic variations of that sequence.

- 27. An amino acid sequence as shown in SEQ ID No. 4, including variants thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.
- 5 28. A cDNA sequence identified herein as SEQ ID No. 5 including non-critical allelic variations of that sequence.
- 29. An amino acid sequence as shown in SEQ ID No. 6, including variants thereof having non-critical amino acid substitution(s) or deletion(s) at one or more locations in that sequence.
  - 30. An AGP enzyme or sub-unit thereof whose *in vivo* activity is substantially independent of any level of *in vivo* 3-phospho-glycerate and/or any *in vivo* level of inorganic phosphate, and whose activity is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.
    - 31. Plasmid pPPS1.
    - 32. Plasmid pPPL1.

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- 33. Plasmid pPPL1M.
- 34. Plasmid pPPS4.
- 25 35. Plasmid pPPL4.
  - 36. Plasmid pPPL5.
  - 37. Plasmid pBKL4.

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38. Plasmid pVictor IV GN.

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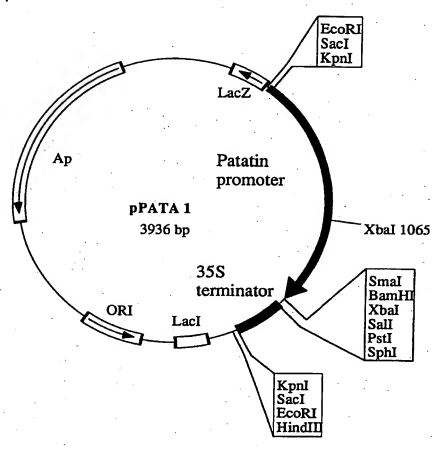
- 39. A method of increasing the rate and/or yield of starch production in an organism, especially a plant or a plant cell, which method comprises introducing into an organism a recombinant DNA construct containing an exogenous DNA sequence encoding an exogenous ADP glucose pyrophosphorylase enzyme (AGP) or sub-unit thereof and one or more promoter sequences enabling the expression of the AGP encoded by that sequence by the organism thereby to increase the AGP content of the organism and in consequence to increase the rate of starch production by the organism and/or the starch yield, characterised in that the said DNA sequence is the gene sequence encoding the barley (Hordeum vulgare) endosperm AGP or a sub-unit thereof, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units, wherein the construct is expressed in the organism; characterised in that the activity of the enzyme or sub-unit thereof is substantially independent of any level of in vivo 3-phospho-glycerate and/or any in vivo level of inorganic phosphate; and further characterised in that the activity of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.
- A vector for the transformation of an organism, especially a plant or a plant 40. cell, to increase the AGP content of such an organism consequently to increase the 20 rate of starch production by such an organism, such vector comprising a recombinant DNA construct containing a DNA sequence encoding an exogenous ADP glucose pyrophosphorylase enzyme (AGP), such vector also incorporating the necessary promoter and other sequences enabling the expression of that exogenous AGP in an organism transformed by that vector, characterised in that the said DNA sequence is the gene sequence encoding the barley (Hordeum vulgare) endosperm AGP or a subunit thereof, or a variant thereof having non-critical amino acid substitution(s) or deletion(s) at one or more points in the amino acid sequences defining the barley endosperm AGP or either of its sub-units, wherein the construct is capable of being expressed in the organism; characterised in that the activity of the enzyme or sub-unit thereof is substantially independent of any level of in vivo 3-phospho-glycerate and/or any in vivo level of inorganic phosphate; and further characterised in that the activity

of the enzyme or sub-unit thereof is not stimulated by fructose-1,6-bisP and/or is not inhibited by AMP.

- 41. A method of targeting an exogenous protein to the amyloplast of plants or plant cells which comprises introducing into the plant or plant cell a recombinant DNA construct containing a DNA sequence encoding a starch branching enzyme transit peptide and an exogenous DNA sequence encoding the exogenous protein; wherein the construct is capable of being expressed in the plant or plant cells.
- 42. A method according to claim 41, characterised in that said DNA sequence encoding the starch branching enzyme comprises the sequence identified as SEQ.ID.No.5.
- 43. A method according to claim 41, characterised in that said starch branching enzyme expressed in the plant or plant cell by said construct comprises the amino acid sequence identified as SEQ.ID. No. 6.
  - 44. A foodstuff made from or comprising an organism according to claim 1 or any claim dependent thereon.
  - 45. A foodstuff according to claim 44 wherein the foodstuff is a fried foodstuff.
  - 46. A foodstuff according to claim 44 or claim 45 wherein the foodstuff is a potato.

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FIG. 1



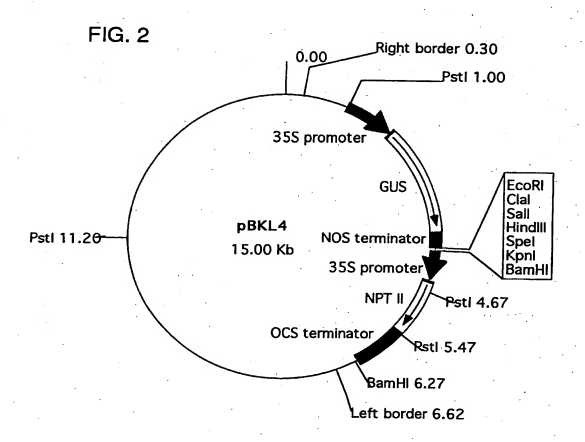
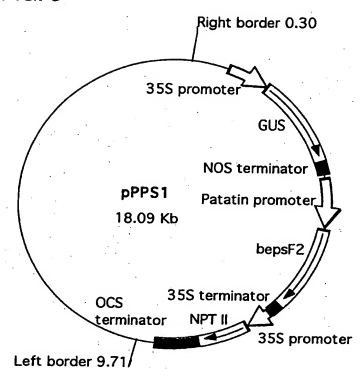


FIG. 3



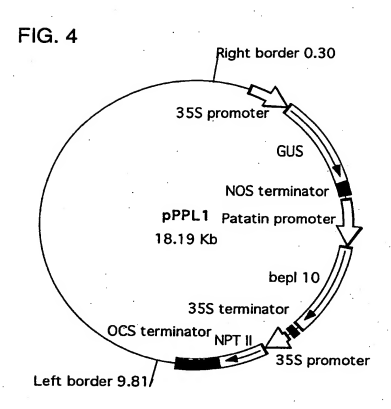
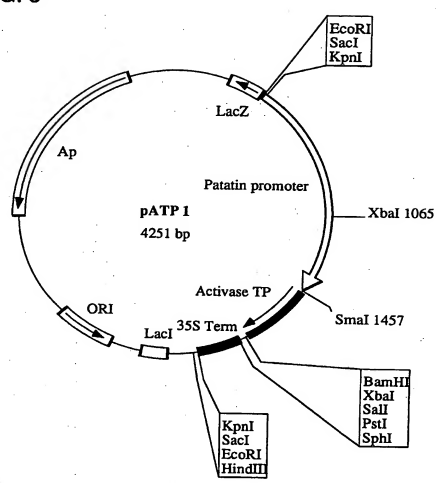
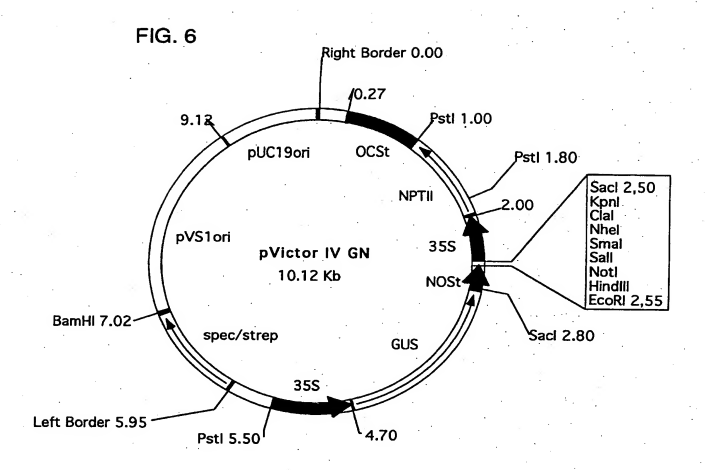


FIG. 5





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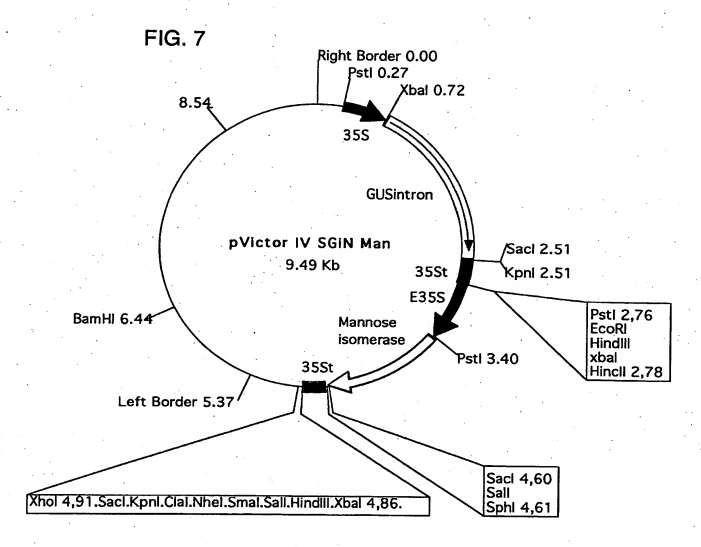
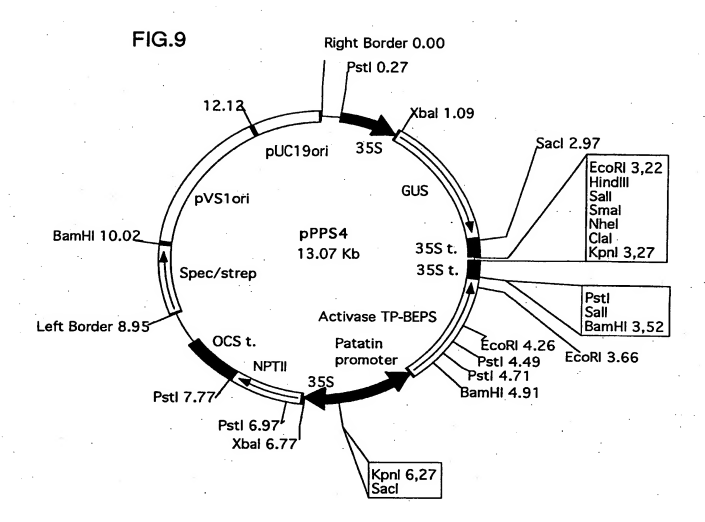
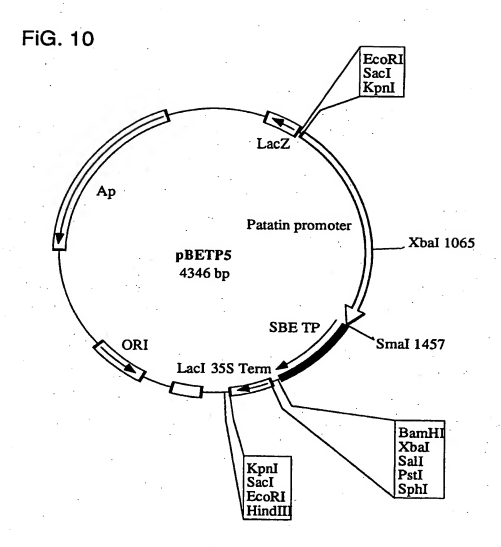
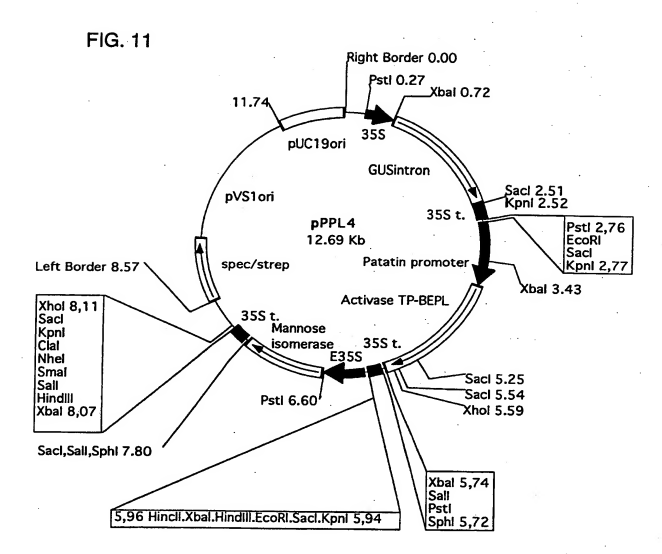


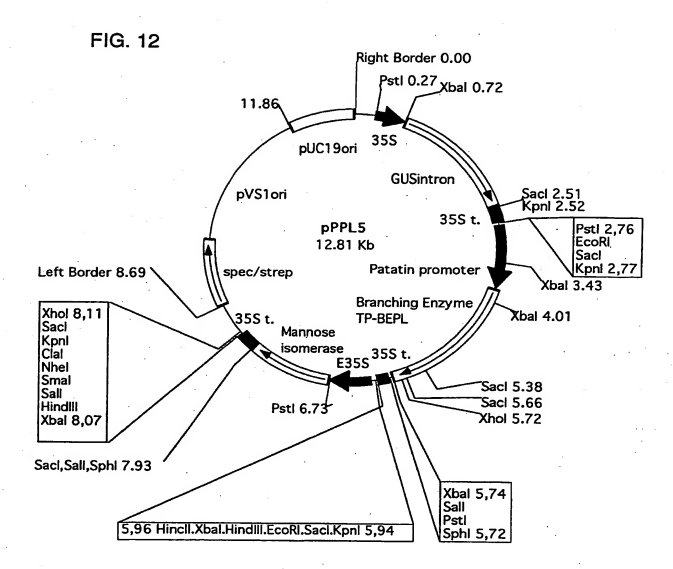
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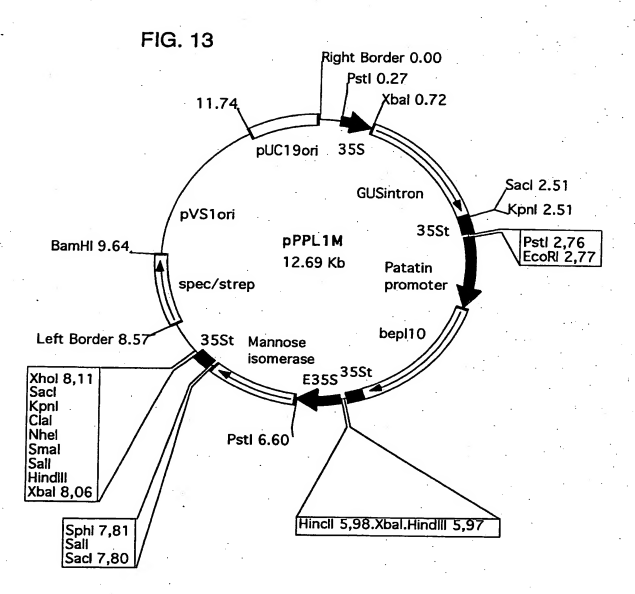
- 8.1 Amino terminal amino acid sequence of the rubisco activase AGP small subunit fusion enzyme
- 1 MATAVSTVGA ATRAPLNLNG SSAGASVPTS GFLGSSLKKH 40
- 41 TNVRFPSSSR TTSMTVKAAE NEEKNTDKWA HLAKDFSDDQ 80
- 81 LDIRRGKGMV DSLGSMDVPL ASKVPLPSPS KHEQCNVYSH 120
- 8.2 Amino terminal amino acid sequence of the rubisco activase AGP large subunit fusion enzyme
- 1 MATAVSTVGA ATRAPLNLNG SSAGASVPTS GFLGSSLKKH 40
- 41 TNVRFPSSSR TTSMTVKAAE NEEKNTDKWA HLAKDFSDDQ 80
- 81 LDIRRGKGMV DSLGIHMQFS SVLPLEGKAC VSPVRREGSA 120
- 8.3 Amino terminal amino acid sequence of the starch branching enzyme AGP large subunit fusion enzyme
- 1 MEINFKVLSK PIRGSFPSFS PKVSSGASRN KICFPSQHST 40
- 41 GLKFGSQERS WDISSTPKSR VRKDERMKHS SAISAVLTDD 80
- 81 NSTMAPLEED VKTENIGLLN LDPMQFSSVL PLEGKACVSP 120











# FIG. 14

Nucleotide sequence and derived amino acid sequence of a cDNA encoding the large subunit of ADP-glucose pyrophosphorylase from barley seed endosperm.

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14b/17

1150 1170 1190 TOCTGAACCTTCTAAAGTCAAGATACGCAGAACTACATGACTTTGGGTCTGAAATCCTCC LNLLKSRYAELHDFGSEILP 1210 1230 CCAGAGCICIGCATGATCACAATGTACAGGCATATGTCTTCACTGACTACTGGGAGGACA RALHDHNVQAYVFTDYWEDI 1270 1290 1310 TTGGAACAATCAGATCCTTCTTCGATGCGAACATGGCCCTCTGCGAACAGCCTCCAAAGT G T I R S F F D A N M A L C E Q P P K F 1330 1350 1370 TIGAATITTATGATOCAAAAACCCCCTTCTTCACTTCGCCTCGGTACTTACCGCCAACAA EFYDPKTPFTSPRYLPPTK 1390 1410 AGTCAGACAAGTGCAGGATCAAAGAAGCGATCATTTCGCACGGCTGCTTCTTGCGTGAAT SDKCRIKEAIISHGCFLREC 1450 1470 1490 GCAAAATCGAGCACTCCATCATCGGCGTTCGTTCACGCCTAAACTCCGGAAGCGAGCTCA K I E H S I I G V R S R L N S G S E L K 1530 1510 1550 AGAACGCCATGATGATGGGCGCGGACTCGTACGAGACCGAGGACGAGATCTCGAGGCTGA NAMMMGADSYETEDEISRLM 1570 1590 1610 TGTCTGAGGGCAAGGTTCCCCATCGGCGTCGGGGAGAACACAAAGATCAGCAACTGCATCA SEGKVPIGVGENTKISNCII 1630 1650 TOGACATGAACGCGAGGATAGGAAGGGACGTGGTCATCTCAAACAAGGAGGGGGTGCAAG D M N A R I G R D V V I S N K E G V Q E 1690 1710 1730 AAGCCGACAGGCCGGAGGAAGGGTACTACATCAGGTCCGGGATCGTGGTGATCCAGAAGA ADRPEEGYYIRSGIVVIQKN

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14c/17

ACGCCACCATCAACCACCGCCACCGTCGTGTACGCCGTCCCCGGTCCGCCCACCGCGTTCC ATIKDGTVV\* TGCCACAACCIGIGCGCIGCGICGGICGTCATCATCTTCTCAAACTCCGGCACTGAAGAA GICATOCGGGGACGCGAGACGITTGAAGCITGAATCACTGACACTGAAAGTGAAGGCGCA 

# FIG. 15

Nucleotide sequence and derived amino acid sequence of a cDNA encoding the small subunit of ADP-glucose pyrophosphorylase from barley seed endosperm.

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A	D	A	V	R	Q	Y	L	W	L	F	E	E	Н	N	V	M				

#### 15a/17

550 570 590 AATTCTTGCTGGAGATCACCTGTACCGAATGGACTATGAAAAGTTTATTCAGGCACACAG ILAGDHLYRMDYEKFIQAHR 610 630 ACAAACGCATGCTCATATTACTGTTGCTCCCTTGCCCATGGATGAGCAACGTGCAACTGC ETDADITVAALPMDEERATA 670 690 ATTTGGCCTTATGAAAATCGATGAAGAAGGGAGGATAATTGAATTCGCAGAGAAACCAAA FGLMKIDEEGRIIEFAEKPK 730 750 770 AGGAGAACAGTTGAAAGCTATGATGGTTGATACGACCATACTTGGCCTTGAAGATGCCAG G E Q L K A M M V D T T I L G L E D A R 790 810 830 GGCAAAGGAAATGCCTTATATTGCTAGCATGGGTATCTATGTTATTAGCAAACATGTGAT AKEMPYIASMGIYVISKHVM 850 870 GCTTCAGCTTCTCCGTGAGCAATTTCCTGGAGCTAATGACTTCGGAAGTGAAGTTAT.CC LQLLREQFPGANDFGSEVXP 910 930 950 TGGTGCAACTAGCACTGGCATGAGGGTACAAGCATACCTATACGACGGTTACTGGCAAGA GATSTGMRVQAYLYDGYWED 970 990 1010 TATIGGIACAATIGAGGCATICIATAATGCAAATITGGGAATIACCAAAAAACCAATACC IGTIEAFYNANLGITKKPIP 1030 1050 1070 TGATTTCAGTTTCTATGACCGTTCTGCTCCCATTTACACACAACCTCGACACTTGCCTCC D F S F Y D R S A P I Y T Q P R H L P P 1090 1110 TTCAAAGGITCITGATGCTGATGIGACAGACAGIGTAATTGGTGAAGGATGTGTTATTAA SKVLDADVTDSVIGEGCVIK

#### 15b/17

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	15	10						153	30						155	0			
AAGAC	AGG	GC.1	IAC.	TIC	GI	CAG	ICIO	GAZ	AIC	AAC	CAA	CAAC	333	330	GAA	• 3GA	TAE	ZATZ	· AA
	157	0			•		1	1590	)		•			1	610			•	
AATAA	AAA	.œ	AG1	rGCC	TE	3003	AGT(	'AC'I	TC:	IAC	400	· CIT	rτα	$\infty$	œr.	IGA:	GIZ	YTT	AG
	16	30						165	50						1670	)			
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GGCACZ	***	<b>7.7.</b> 7	·V·V·	HHA	HAA	MAA	٠									•			

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-	LJ.	- 1	n

Nucleotide sequence and derived amino acid sequence of a cDNA encoding a potato starch branching enzyme.

60

CCCGTCTGTAACCATCATTAGTCATGTTGTTCCACCTCAATGCCATCATTCACATCCAAA

120

CGTCTGGGGTGAGAACATACAAGAAGGCAGCAGCTGAAGCAAAGTACCATAATTTAATCA

180

ATGGAAATTAATTTCAAAGTTTTATCAAAACCCATTCCAGGATCTTTTCCCATCTTTCICA M E I N F K V L S K P I R G S F P S F S

240

CCTAAAGTTTCTTCAGGGCCTTCTAGAAATAAGATATGTTTTCCTTCTCAACATAGTACT P K V S S G A S R N K I C F P S Q H S T

300

GCACTGAAGTTTGGATCTCAGGAACGGTCTTGGGATATTTTCTTCCACCCCAAAATCAAGAG L K F G S Q E R S W D I S S T P K S R

360

GITAGAAAGATGAAAGGATGAAGCACAGTTCAGCTATTTCCGCTGTTTTGACCGATGAC V R K D E R M K H S S A I S A V L T D D

420

AATTOGACAATGGCACCCCTAGAGGAAGATGTCAAGACTGAAAATATTGGCCTCCTAAAT N S T M A P L E E D V K T E N I G L L N

**4**20

TIGGATCCAACTITIGGAACCTTATCTAGATCACTTCAGACACACACACACTATGTG L D P T L E P Y L D H F R H R M K R Y V

#### FIG. 17

Nucleotide sequence of a tuber specific class I patatin promoter.

# 17a/17

660
ACICICATTICAACICAGITTAAGCAATTCICATAAGGCCAGCAAAATCACAGIGCTGAA
720 ATCIAGAAAAATCICATACAGIGAGATAAATCICAACAAAAACGITGAGIOCATAGAGAG
ALCTHERANTICICATIONALITY AND
780 GGIGIATGIGACACCCAACCTCAGCAAAAGAAAACCTCCCCTCAAGAAGGACATTTGCGG
840 TCCTAAACAATTTCAAGTCTCATCACACATATTATTATTATTATTAATTA
900 AAAAGGAAAGTGAGAGATAATGACAGTTGCGGTGCAAAGTGAGTG
960 ATCAGIAATAGACATCACIAACITTTATTGGITATGICTTTCTCAAAATAAAAT
1020 ACTIGITIACGIGCCIATATATACCATGCTTGTTATATGCTCAAAGCACCAACAAAATTT
1047
AAAAACACTITGAACATTIGCCCCCGG

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